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(57) Abstract <p>DNA molecules encoding a family of zinc-finger DNA binding domains, which appears to function to monitor levels of a superoxide-dependent signal and negatively regulates a plant cell death pathway, including wild-type LSD1, LOL1 and LOL2, and proteins which physically interact with LSD1, indicating a function with LSD1 of controlling plant cells' response to pathogens.</p>			

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PLANT PATHOGEN RESPONSE GENE

CROSS-REFERENCE TO RELATED APPLICATIONS

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This application claims benefit of U.S. Provisional Application No. 60/039,063 filed February 28, 1997.

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BACKGROUND OF THE INVENTION

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Field of the Invention

This invention relates to a novel DNA molecule that encodes a novel polypeptide, LSD1, which has an effect in regulating the initial response of plants to pathogens and the subsequent spread of plant cell death engendered by infection, the protein encoded by the gene, and transgenic plants comprising the DNA molecule. This invention also relates to novel DNA molecules encoding LSD1 related proteins LOL1 and LOL2. In addition, it relates to novel DNA molecules encoding proteins which directly interact with LSD1.

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Description of the Related Art

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Controlled induction of cell death occurs during both normal plant development and as the rapid, localized response to pathogen infection known as the hypersensitive response (HR) (Stakman, 1915; Goodman and Novacky, 1994; Dangl et al., 1996). The HR is a feature of most, but not all, disease resistance reactions in plants. The disclosure of these publications and all others cited herein, as well as of the priority application, is incorporated herein by reference.

30

Genetic control of disease resistance reactions is of two broad classes. The first is determined by specific interactions between particular alleles of pathogen *avr* (avirulence) gene loci and an allele of the corresponding plant disease resistance (*R*) locus. When these alleles are present in both host and pathogen, the result is disease resistance in the plant, and the interaction is said to be "incompatible". If either the plant *R* allele or the cognate pathogen *avr* gene are absent or inactive, disease results and the interaction is said to be "compatible" (reviewed by Flor, 1971; Crute, 1985; Keen, 1990; Pryor and Ellis, 1993). A great deal of progress has been made recently in understanding the molecular structure of *R* genes and their predicted products (reviewed by Dangl, 1995; Staskawicz et al., 1995; Bent, 1996). These molecules function to recognize *avr* dependent signals and trigger the plant cell to begin the chain of signal transduction events culminating in a halt of pathogen growth. The simplest mechanistic interpretation of allele-specific disease resistance is that the *R* gene product recognizes the *avr* gene product directly. Although no direct *avr-R*

protein interaction has been shown in planta, expression of *avr* genes in plant cells can be sufficient to trigger the HR in a *R*-dependent manner, and *avr-R* protein-protein interactions can occur in yeast two-hybrid systems (Gopalan et al., 1996; Scofield et al., 1996; Tang et al., 1996).

5 The second mode of genetic control of disease resistance is termed "non-host" resistance and describes in essence those interactions which lack genetic variability in either host or pathogen such that no virulent pathogen and no susceptible host line have been identified. While it is not beyond reason to assume that traditional "non-host" interactions are simply a series of allele specific recognition events occurring simultaneously (Whalen et al., 1988; Kobayashi et al., 1989; Valent et al., 1990), it is also possible that this mode of 10 resistance is mechanistically distinct from that mediated by allele-specific interactions. Pathogen ligands (termed elicitors) which mediate several key non-host interactions have been isolated, although their corresponding plant receptors have not (Cosio, et al. 1992; Nürnberg et al., 1994).

15 Subsequent to pathogen recognition by either of these two systems, the plant cell deploys a battery of inducible defense responses. Chief among the earliest events are calcium influx, $K^+ - H^+$ exchange leading to alkalinization of the extracellular space, and an oxidative burst (reviewed in Godiard et al., 1994; Hammond-Kosack and Jones, 1996). The latter is potentially mediated by a plasma membrane NADPH oxidase analogous to that used by mammalian neutrophils (Low and Merida, 1996), although other models exist 20 (Bolwell et al., 1995). Parts of this cascade are linearly regulated in at least some systems: blocking of Ca^{2+} influx blocks anion channel activity, the oxidative burst and downstream events including cell death; blocking anion channels effects only ROI production and defense gene activation, but not Ca^{2+} influx (Nürnberg et al., 1994; Levine et al., 1996; 25 May et al., 1996).

Consequent production of reactive oxygen intermediates (ROI) occurs with kinetics and magnitude suggesting a key role in either pathogen elimination, subsequent signaling of downstream effector functions, or both (reviewed by Baker and Orlando, 1995; Low and Merida, 1996). H_2O_2 can have a key role in resistance responses, and cell wall 30 strengthening (Brisson et al., 1994; Levine et al., 1994; Levine et al., 1996), and superoxide produced as the proximal ROI in the burst has also been implicated in initiating HR (Doke, 1983; Jabs et al., 1996). Transcription and translation of plant genes are required for HR. These signals are thought to culminate in transcriptional activation of a variety of plant 35 genes, HR, and the production of both local and systemic signals that protect the plant from further infection. It is unclear whether these effector functions are controlled by linear, interdigitating, or bifurcating signal pathways.

Cell death during the HR may be a direct consequence of ROI toxicity, or it may be

a secondary consequence of signals derived from ROI. It is not known whether HR is required to halt pathogen growth. Nonetheless, HR is correlated with the onset of systemic acquired resistance (SAR) to secondary infection in distal tissue (reviewed by Ryals et al., 1996). In at least tobacco and Arabidopsis, enzymatic blocking of salicylic acid (SA) accumulation subsequent to infection alters disease resistance responses, and SA in distal tissues is required for SAR (Gaffney et al., 1993; Delaney et al., 1994; Vernooij et al., 1994). SA accumulates following the oxidative burst to high levels locally at infection sites. The biochemical properties of SA as an inhibitor of a variety of enzymes suggest a model whereby SA or a radical derived from it poisons the infected cell, causing its death (Enyedi et al., 1992; Malamy et al., 1992; Chen et al., 1994; Durner and Klessig, 1995; Rueffler et al., 1995). Recent descriptions of the morphology of cell death during infection suggest, in at least some cases, parallels with animal apoptosis (Mittler et al., 1995; Kossak et al., 1996; Levine et al., 1996; Ryerson and Heath, 1996; Wang et al., 1996a; reviewed by Dangl et al., 1996). A molecular understanding of both the signaling events that control the onset of this specialized plant cell death and the mechanisms by which these cells die will hasten approaches to manipulate cell death to protect plants from disease.

A number of researchers have isolated mutants in Arabidopsis which exhibit constitutive onset of HR-like cell death in the absence of pathogen (Greenberg and Ausubel, 1993; Dietrich et al., 1994; Greenberg et al., 1994). These mutants resemble a variety of mutants in crop species isolated since the 1920s and broadly categorized as "lesion mimic mutations" (Langford, 1948; Kiyosawa, 1970; Walbot et al., 1983; Johal et al., 1994). A series of non-allelic mutations was isolated which expressed histochemical and molecular markers associated with disease resistance responses. These mutants subdivide the lesion mimic class into a "lesions simulating disease resistance" or *lsd* phenotype (Dietrich et al., 1994). These mutants also exhibited heightened resistance to otherwise virulent bacterial and oomycete pathogens when lesions were present, demonstrating that these cell death phenotypes can trigger pathogen non-specific resistance resembling SAR. Similar "accelerated cell death" or *acd* mutants have been described by Greenberg and Ausubel (Greenberg et al., 1994). Greenberg and Ausubel (1993) additionally isolated a mutant which though expressing an *acd* phenotype was in fact more susceptible to pathogen. It is thus possible to identify genetically at least two types of cell death, namely those which feed into a pathway culminating in establishment of a disease resistant state, and those which do not.

The *lsd1* mutant is exceptional. In conditions permissive for wild type plant growth and in the absence of detectable microscopic lesions, the *lsd1* mutant is hyper-responsive to challenge by a variety of stimuli including pathogens and low doses of chemicals which trigger the onset of SAR (Dietrich et al., 1994). Mutant *lsd1* plants are resistant to

otherwise virulent pathogens in conditions where no spontaneous cell death lesions form. Following initiation of cell death in a local spot on a leaf, lesions propagate throughout the leaf and kill it 2-4 days later. Propagation of locally initiated cell death is confined to the inoculated leaf. Thus, *LSD1* functions to negatively regulate both the initial response to pathogens and the subsequent spread of cell death. Superoxide is a necessary and sufficient trigger for this phenotype, and superoxide production precedes onset of cell death by 8-16 hours following initiation by three different triggers (Jabs et al., 1996). Therefore, the *LSD1* gene responds to either superoxide or to a signal derived from it to down regulate or dampen the cell death response, resulting in the typical locally bounded HR. The invention herein includes the *LSD1* gene, which encodes the first member of a new subclass of zinc-finger proteins in *Arabidopsis*.

It is therefore an object of the invention to provide a novel DNA molecule, *LSD1*, isolated from *Arabidopsis* which works to protect plant cells in response to pathogens, and DNA molecules encoding *LSD1* related proteins *LOL1* and *LOL2*.

It is a further object of the invention to provide the protein encoded by *LSD1*, and transgenic plants comprising *LSD1*. Knowledge of the structure of the *LSD1* gene allows accurate creation of particular mutants (e.g., deletion and point mutations), for example, mutants having a dominant negative phenotype, analogous to the mutants of *Drosophila PANNER* gene (Ramain et al., 1993), using methods known in the art. This in turn allows engineering of transgenic crop plants which do not suffer cell death, but are still resistant to infection. In addition, expression of the dominant negative *LSD1* protein may be refined so that it is expressed very quickly after infection.

The *LSD1* protein is also a useful target for herbicide development. Transgenic plants may be made in which *LSD1* mutant genes are expressed which are resistant to herbicidal compounds which normally result in cell death in combination with the wild-type *LSD1*. Mutants of the *LSD1* gene are tested in a *lsl1* background to determine if the mutant has a normal or novel function, and in a wild-type background to determine the existence of a dominant negative function.

Other objects and advantages will be more fully apparent from the following disclosure and appended claims.

SUMMARY OF THE INVENTION

The invention herein comprises the DNA molecule of the wild-type *LSD1*, which functions to monitor levels of a superoxide-dependent signal and negatively regulates a plant cell death pathway. The predicted *LSD1* protein contains three zinc-finger domains, defined by CxxCxRxxLMYxxGASxVxCxxC (SEQ ID NO:54). The invention further comprises a protein encoded by *LSD1*, and transgenic plants comprising *LSD1*, and

mutations thereof.

In particular, the preferred embodiments of the invention herein include the following: an isolated DNA molecule, encoding the LSD1 polypeptide sequence, selected from the group consisting of SEQ ID NOS:13-15; the LSD1 DNA molecule having the nucleotide sequence as set forth in SEQ ID NO:13; the DNA molecule that is cDNA; the DNA molecule which is genomic DNA; a chimeric construction comprising a promoter sequence and the LSD1 DNA molecule or portions of the LSD1 DNA molecule; a recombinant plant transformed with the LSD1 DNA molecule; a transformed plant comprising a DNA molecule encoding a protein as set out in SEQ ID NO:16 or SEQ ID NO:17; an isolated protein molecule comprising the protein set out in SEQ ID NO:16 or SEQ ID NO:17; a transformation vector comprising a LSD1 DNA molecule as set forth herein; an isolated DNA molecule encoding the zinc finger consensus sequence shown in SEQ ID NOS: 1-3; and anything that hybridizes to the LSD1 DNA molecule set forth herein under hybridization conditions as defined herein.

Other objects and features of the inventions will be more fully apparent from the following disclosure and appended claims.

BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A-C show the physical delineation of the *lsd1* mutation. Figure 1A shows YAC clones at *lsd1*. The arrowheads imply the YAC clone extending in the direction given, solid vertical black bars denote YAC ends used to isolate genomic phage clones and subsequently converted into CAPS RFLP markers as described (refer to Figure 2 for their map position and to Tables 1 and 2, Examples II and III, for their definition). Figure 1B shows the three BAC clones which contained the CAPS markers listed above BAC1G5. The arrowheads imply extension of the BAC clone in the direction shown. The scale in Figures 1A and 1B are the same. Figure 1C shows the genomic phage clones positioned under an expansion of three of the BACs. The diamond-filled bar represents the 8A6-1.3 clone, which co-segregated with *lsd1*, used to isolate these phage. The *lsd1* deletion is noted at the bottom.

Figure 2 is a genetic linkage map of the *lsd1* region. The vertical line at the left represents the section of *Arabidopsis* chromosome 4 between CH42 and B9-1.8 (telomeric toward bottom). CAPS-based RFLP markers discussed in the text intersect the chromosome, and their relative recombination frequencies in the F₂ mapping population are placed in the center. The number of meioses identified among the total number of F₂'s scored is at the right. The arrowhead denotes the co-segregating marker.

Figures 3A-C show molecular fine mapping of the *lsd1* locus. Figures 3A and 3B show genomic DNA blots demonstrating the presence of a 0.8 kb deletion om the *lsd1*

mutant. Genomic DNA (5 g) from wild type Ws-0 or *lsd1* was digested with (for each pair of lanes from left to right) EcoRI, HindIII, a double digest of HindIII and XbaI, or KpnI. In Figure 3A, the blot was probed with the 0.8 kb EcoRI-XbaI. In Figure 3B, a duplicate blot was probed with the 4.5 kb PstI-XbaI fragment. The probes are depicted in Figure 3C, and were isolated from phage clones depicted in Figure 1C. Molecular weight markers are the Gibco-BRL 1 kb ladder. Figure 3C shows the restriction map in and around the *lsd1* gene. The extent of the deletion of this locus is shown as are the extent of the hybridization of the various restriction fragments with *lsd1* cDNAs. Genomic restriction fragments used in complementation experiments are underlined. The asterisk refers to an XbaI site derived from the phage lambda cloning junction.

Figure 4 shows that the *lsd1* mutation is an mRNA null allele. RNA blots (1 g of polyA+ RNA) from leaf tissue of 5 week old plants kept in short days (permissive for *lsd1* growth) 3 days after spraying with either INA (0.3 mg/ml powder containing 25% active ingredient, or 4 mM), or wettable powder control. Spreading *lsd1* lesions had just started to appear at the time of leaf harvest. Probes were purified inserts from the LSD1 cDNA as represented by EST 82D11T7 (top), a PR-1 cDNA (Uknes et al, 1993b), and an actin cDNA. The blot was probed successively in the order displayed.

Figure 5 shows the zinc finger domains (SEQ ID NOS:1-3) of the predicted LSD1 protein and the alignment of the three zinc finger domains. The numbers at the left and right refer to amino acid residue position in the deduced LSD1 protein. Vertical lines indicate conservation in pairwise comparison, and a colon indicates conservative substitution. A consensus sequence is listed below, with conservative substitutions noted in the second line of consensus where "+" is basic, plus charged; and "@" is amide, polar, uncharged, hydrophilic.

Figure 6 shows how the carboxyl portion of the deduced LSD1 protein is related to known DNA-binding and transcription factors. Vertical lines indicate conservation in pairwise comparison, and a colon indicates conservative substitution. Figure 6A shows homology of a slightly longer portion of the deduced LSD1 protein with mammalian insulin receptor substrate proteins. The LSD1 translation product (SEQ ID NO:4) is shown on the top, aligned with the mouse insulin receptor substrate (SEQ ID NO:5). In this region, all mammalian insulin receptor substrates are identical. Figure 6B shows the homology of LSD1, on each top line, with four known transcription factors. The LSD1 translation product (SEQ ID NO:6) is shown on top, and below it are the related domains from a human early growth response (EGR) Zn-finger protein (SEQ ID NO:7, a human TGF--early induced Zn-finger protein (SEQ ID NO:8), a *Xenopus laevis* H-L-H transcription factor (SEQ ID NO:9), and the human ELK-1 protein (SEQ ID NO:10). Figure 6C shows the homology of a LSD1 transcription product (SEQ ID NO:11) with a putative maize

transcription initiator binding protein (SEQ ID NO:12). GenBank accession numbers of each protein are listed at the right.

Figure 7 shows the consensus sequence of the zinc finger domains (SEQ ID NOS:63-65, respectively) of LSD1 (A), LOL1 (B) and LOL2 (C).

5 Figure 8 shows the homologies between the first (A), second (B) and third (C) zinc finger domains of LSD1, LOL1 and LOL2

DETAILED DESCRIPTION OF THE INVENTION AND PREFERRED EMBODIMENTS THEREOF

10 The present invention provides a genomic DNA sequence (SEQ ID NO:13) and a cDNA sequence (SEQ ID NOS:14-15) or the *LSD1* gene which is required for the regulation of initial plant response to pathogens, and cDNA proteins deduced (from short form, MG7-SEQ ID NO:16; from long form, MG, SEQ ID NO:17).

15 In addition, the invention herein provides functional protein domain sequences involved in regulating genes controlling cell death. Gene expression can be regulated by attaching a promoter to the *LSD1* gene, which may be either the native promoter or any other promoter.

20 The invention herein includes the DNA molecule having the nucleotide sequence as set forth in SEQ ID NOS:13, 14 and 15, encoding either of two *LSD1* polypeptides, which are preferably the *LSD1* polypeptides set forth in SEQ ID NOS:16 and 17. This DNA molecule may be cDNA or genomic. The invention also includes as the open reading frame any chimeric construction comprising a promoter sequence and the DNA molecule of the invention, a recombinant plant transformed with the DNA molecule, and any transformation 25 vector comprising the DNA of the invention. In addition, the DNA sequence of either the full-length SEQ ID NO:13, or a shortened or otherwise modified version thereof, may be modified to optimize its expression in plants, with codons chosen for production of the same or a similar protein as encoded by the wild type *LSD1* gene. Other modifications of the *LSD1* gene that yield a protein having essentially the same properties as the *LSD1* gene 30 are included within the invention herein.

The invention herein also includes anything that hybridizes to the *LSD1* DNA (SEQ ID NO:13) of the invention as discussed above, under hybridization conditions, which are defined as: 7% Na dodecyl sulfate (SDS), 0.5 M sodium phosphate, pH 7.0, 1 mM EDTA at 35 50C, and wash in 2X SSC buffer, 1% SDS, at 50C (Church and Gilbert, 1984). Proc. Natl. Acad. Sci. USA 81:1991-1995 (1984)).

The novel *LSD1* gene of the present invention, it its wild type form or as mutated by selected mutations and genetically engineered derivatives obtained as is known in the art,

and proteins encoded thereby, are included in the invention herein, and may be transferred into any plant host using methodology known in the art for purposes of altering the extent and type of plant resistance to pathogens, and to change resistance to particular herbicides.

The mutant phenotype of the null *lsd1* allele suggests that the wild type product is a negative regulator of cell death. In addition, *lsd1* reacts to both nominally virulent pathogens, and to chemicals which trigger the onset SAR, with an HR-like response. But it is important to note that *lsd1* expresses wild type timing of *R* gene driven HR (Dietrich et al., 1994)--it is the subsequent spread of cell death which distinguishes the mutant. Thus, cell autonomous signals required for *R* gene function are intact in an *lsd1* null, but the response to cell non-autonomous signals emanating from cells undergoing HR is perturbed. Collectively, these features of the mutant phenotype suggest that LSD1 functions to limit both the initiation of defense responses and the subsequent extent of the HR. The fact that an *lsd1* null is hyper-responsive to signals initiating the defense response and HR-like cell death additionally suggests that these pathways are functionally intact in the wild type cell, but require a threshold level of signal for full activation.

LSD1 appears to act as a transcription factor (or as a protein which sequesters a transcription factor). As outlined above, the oxidative burst in an infected cell generates a superoxide-dependent signal up-regulating the HR pathway. This signal overcomes the negative regulatory function of the available LSD1, and drives primary responding cells into the HR pathway. Additionally, the cells undergoing HR amplify the signal, probably via a sustained extracellular oxidative burst, to neighboring cells. The primary signal molecule may be diffusible over short ranges (Levine et al., 1994), could act as an autocrine signal, and could lead to the accumulation of a secondary signal molecule in a steep spatial gradient from the infection site. At a critical point in the signal gradient, a threshold is reached. Above that point the pro-death pathway operates, and below it the pro-death response would be attenuated by LSD1. Such a gradient is formed by SA and SA-conjugates (Enyedi et al., 1992); SA biosynthesis can be induced by hydrogen peroxide (Leon et al., 1995); and sub-effective doses of SA can amplify pathogen-derived signals (Kauss et al., 1992; Kauss and Jeblick, 1995; Mauch-Mani and Slusarenko, 1996). Thus, it could be that an SA gradient dictates LSD1 activity.

Constitutive expression levels by LSD1 could suffice to protect cells below the critical signal threshold for death induction. The time lag of 12-16 hours observed between superoxide production initiated in *lsd1* by a variety of triggers and the onset of cell death (Jabs et al., 1996), which could provide sufficient time for up-regulation of LSD1 activity before irrevocable commitment to death during wild type responses, so that cell death could spread until sufficient active LSD1 accumulates. Alternatively, this time lag could represent a requirement for biosynthesis of pro-death intermediates and LSD1 normally could operate

by interdicting this pathway. LSD1 could positively regulate anti-cell death targets, potentially including genes involved in cell survival, ROI de-toxification, or in degradation of a key intermediate in the pro-death pathway. Alternatively, LSD1 could act as a transcriptional repressor directly on genes in the pro-death effector pathway. This scenario 5 differs from the first only in that the set of target genes would be different. The availability of extragenic suppressors of *lsd1* will aid in identifying LSD1 targets (Jabs et al., 1996).

This model also explains the runaway cell death phenotype of the null *lsd1* mutant. In the absence of LSD1, the threshold normally required before commitment to HR is removed. Thus, minimal up-regulation of the superoxide-dependent signal drives the cell 10 into the HR pathway. Hence the ability of *lsd1* to respond to virulent pathogens as if resistant derives from a lack of background inhibition of the HR pathway normally operating in the cell. Moreover, extracellular superoxide produced during the oxidative burst initiates the same series of events in cells immediately surrounding the site of initiation, and the cell death propagation indicative of the *lsd1* phenotype results. Because 15 the null *lsd1* mutant still requires superoxide for initiation of cell death propagation, it is unlikely that superoxide directly regulates LSD1 activity. This further suggests that a superoxide-dependent signal is the autocrine which propagates the response to neighboring cells.

The *A. thaliana lsd1* mutant phenotype is characterized by enhanced disease 20 resistance, spontaneous formation of lesions in the absence of cell death initiators and failure to limit the extent of cell death. The wildtype LSD1 protein therefore negatively regulates a cell death pathway involved in plant defense responses.

The *LSD1* gene encodes a protein containing a novel zinc finger protein, which is included in the invention herein and is defined by its three consensus zinc fingers: 25 CxxCRxxLMYxxGASxRxVxCxxC (SEQ ID NO:52). These three zinc finger domains have not been observed before in the range of zinc finger proteins. As shown in Dietrich et al., 1997, the *LSD1* gene is a key negative regulator of hypersensitive cell death in plants. We sought other versions of this consensus zinc finger sequence in other plant proteins.

The data on homologies between the LSD1 and LOL1 and LOL2 zinc finger 30 domains indicates that LSD1 as well as LOL1 and LOL2 are members of a novel subclass of zinc finger proteins that are involved in plant cell death pathways. LOL1 and LOL2 might function in cell death phenomena leading to hypersensitive response and disease resistance as has been shown for LSD1. The homologues may also be involved in programmed cell death (PCD) pathways occurring in plants. Examples of PCD in plants include lateral root development, tracheary element differentiation, and abscission of leaves. Preliminary expression studies suggest that LOL2 is expressed in flowers and siliques. Thus a role for LOL2 in PCD pathways leading to petal senescence, anther dihiscence or

PCD of nucellar cells is not unlikely. It is also possible that LOL2 is involved in the hypersensitive response and disease resistance in flowers, thus protecting seeds and ultimately the following generations from pathogen. Alternatively, LOL2 could be up-regulated during the hypersensitive response. Use of *LOL1* and *LOL2* should allow prediction of the protein's function with respect to protection from programmed cell death.

The consensus sequences defined by the LSD1, LOL1 and LOL2 zinc finger domains (Figures 7-8) are thus far unique in the available deduced protein databases. Because zinc finger domains of this type bind DNA and thereby regulate gene activation, it is highly likely that the consensus zinc finger domain defined here is required for proper regulation of related sets of genes. Furthermore, because zinc finger DNA binding domains of related sequence generally control related cellular processes, the new consensus defined here should also do so. Because LSD1 is known to negatively regulate cell death induced by pathogens, it is highly likely that LOL1 and LOL2 also control plant cell death. Thus, the utility of this portion of the invention lies in production of transgenic plants which have mutated versions of the *LOL1* or *LOL2* genes or which overexpress these proteins. Such plants will likely be more resistant to pathogen attack, if, in the first case, the *LOL* genes function to repress defense response (as does *LSD1*). Alternatively, if the *LOL* genes function to activate defense mechanisms, then overexpression will lead to a more effective pathogen response. Because zinc finger proteins featuring other non-LSD1 type DNA binding domains function to either activate or repress gene transcription, we cannot distinguish at present between these two models.

The invention also includes plant proteins, and the genes which encode them, which directly interact with LSD1 protein. Gene regulation in response to pathogen attack is controlled, in part, by the repression and activation of genes. The LSD1, LOL1 and LOL2 proteins encode a novel branch of the zinc-finger DNA binding protein superfamily with roles in controlling plant cell death. As such, they are expected to interact with other proteins. Paradigms of gene activation currently demonstrate that DNA binding proteins can have two classes of "partners". The first class sequesters the DNA binding protein in the cell's cytosol. These partner proteins hold the DNA binding protein out of the nucleus until the correct cellular stimulus is received. This stimulus disrupts the physical interaction, and the DNA binding protein is free to migrate into the nucleus and activate or repress transcription. The second class of protein which interacts with DNA binding protein is made up of proteins which are partners having the role of "enhancing" the gene activation or repression encoded by the DNA binding protein. These partners are termed "co-activators" or "co-repressors" and they may or may not have intrinsic DNA binding activity. We have identified several genes whose protein products interact physically with the LSD1 protein using a common assay, called a "yeast two-hybrid interaction trap" to detect such

interactions genetically (Fields and Sternglanz, 1994; Finley and Brent, 1996). Because the inactivation of LSD1 by mutation leads to enhanced disease resistance, the LSD1 partner proteins represent novel targets for engineering plants with enhanced resistance to pathogens. Thus, this invention includes all proteins which interact with the cell death regulator LSD1 (SEQ ID NOS: 66-91 (includes sequential pairs of nucleic acids and corresponding amino acid sequences).

The features of the present invention will be more clearly understood by reference to the following examples, which are not to be construed as limiting the invention.

EXAMPLES

Example I Care and maintenance of plants

Plants were grown in a chamber at 9 hours light per day, 22°C day temperature and 20°C night temperature essentially as described (Dietrich et al., 1994).

Example II Isolation of DNA and RNA, probe preparation, cloning

Small scale genomic DNA preps were made from single leaves (~1cm long rosette leaves) (Lukowitz et al., 1996). The DNA pellet was re-suspended in 50 ml of Tris/EDTA (TE) and 1 ml was used in a 20 ml polymerase chain reaction (PCR). Large scale genomic DNA preps were done based on the protocol of (Rogers and Bendich, 1985), modified such that concentration in the 2X hexadecyltrimethylammonium bromide (CTAB)(Sigma, St. Louis, MO) buffer was increased to 3% and the precipitated DNA was resuspended in Tris/EDTA/sodium chloride (TEN) buffer and digested with 100 mg/ml, followed by two extractions with chloroform/iso-amyl alcohol and a final precipitation.

RNA was isolated by grinding fresh tissue in liquid nitrogen to a fine powder and extraction in 1 ml of Trizol reagent (Gibco-BRL, Gaithersburg, MD) per 100 mg tissue fresh weight. RNA was isolated according to the manufacturer's protocol. PolyA+ RNA was isolated using DynaBeads (Dynal, Oslo, Norway). RNA blots were formaldehyde agarose gels and contained either 15 mg total RNA or 1 mg polyA+ RNA. HyBond filters for DNA or RNA blots (Amersham, Little Chalfont, United Kingdom) were hybridized in 6xSSC, 5X Denhardt's solution, 0.1% SDS and 100 mg/ml sheared Herring sperm DNA at 65°C. Washes were in 0.2X SSC, 0.1% SDS at the same temperature. RNA blots were stripped for re-hybridization in 5 mM TRIS/2mM EDTA, (pH8.0), 0.1X Denhardt's solution for 1 hour at 65°C.

Example III Isolation of new CAPS markers and genetic mapping of *lsd1*

After establishing linkage to the agamous (*AG*) co-dominant amplified polymorphic sequences (CAPS) marker (Konieczny and Ausubel, 1993), we subcloned and end-sequenced a 1.6 kb HindIII fragment from the RFLP cosmid marker g3883 (position 73.5 on the *Arabidopsis* RI map; Lister and Dean, 1993; see http://nasc.nott.ac.uk/RI_data/top_frame.html), and primers designed based on this sequence. This primer set amplified a rapid amplified polymorphic DNA (RAPD) marker (size difference in Ws-0 versus Col-0 without restriction digestion), and map data generated using this primer allowed us to place *lsd1* below (telomeric to) it. Probe B9-1.8, isolated as a 1.8 kb SstI-EcoRI fragment from the JGB9 genomic phage clone (RI map position ~75; gift of Dr. George Coupland, Cambridge Laboratories, Norwich U.K.) was converted into a CAPS marker. Mapping of this polymorphism placed *lsd1* above (centromeric to) it (Fig. 2). Recombinants were identified as homozygous for one of these CAPS markers, and heterozygous for the other using DNA from F2 individuals. F3 progeny from these recombinants were then scored as either homozygous *lsd1*, segregating *lsd1*, or homozygous wild-type for lesion spread. All CAPS markers we developed are described in Table 1 (below).

Table 1. New PCR based RFLP (CAPS) markers derived during cloning of *lsd1*

<u>Marker</u>	<u>Enzyme</u>	<u>PCR prod.</u>	<u>Col-0</u>	<u>Ws-0</u>
ch42	Clal	1.4 kb	750 bp 650	1.4 kb
g3883-1.6	none		1.4 kb (uncut)	0.7 kb (uncut)
g13838-1.4	HinfI	1.4 kb	450 bp 330 280 200	450 bp 330 280 160
B9-1.8	HinfI	1.8 kb	420 bp 260 240 180 160 140	420 bp 260 240 180 160 140
1H1L-1.6	Ddel	1.6 kb	1.0 kb 300 bp	700 bp 300 (doublet?)
5F7R-1.5	NlaIV	1.5 kb	1.0 kb	1.2 kb

13

			250 bp	250 bp
			200 bp	
20B4-1.6	Ddel	1.6 kb	900 bp	700 bp
			400	400
5				220
			180	180
8A6-1.3	TaqI	1.3 kb	800 bp	800 bp
			400	250
			220	150

10

Example IV Map refinement

YACs were defined (Schmidt et al., 1995; Schmidt et al., 1996, <http://genome-www.stanford.edu/Arabidopsis/JIC-contigs.html>), confirmed by DNA blotting to establish a contig and their ends were isolated by vectorette PCR as described (Matallana et al., 1992; Grant et al., 1995). These ends were also used to isolate genomic phage from a Ws-0 genomic library (Fig. 1). Insert fragments of 1-3 kb were cloned into PBS and end sequenced for derivation of primers identifying new CAPS. PCR conditions (DNA Engine MJ Research) for all CAPS primer pairs except 8A6-1.3 and *lsd1* deletion primers are: 92°C, 3'; 35 cycles of (denature 92°C, 30"; anneal 50°C, 30"; extend 72°C, 2'30"); 72°C, 3'. For 8A6-1.3 and the *lsd1* deletion primer pairs we used 53°C annealing. Table 2 shows the primer sequences used to identify new CAPS markers.

Table 2. Primer sequences used to identify new CAPS markers used for cloning *lsd1*

25	ch42 for	5'-cag tgg atc ttt cct cag acg-3' (SEQ ID NO:18)
	ch42 rev	5'-cat ctt ctt ctg caa tct ggg-3' (SEQ ID NO:19)
	g3883-1.6 for	5'-cat cca tca aac aaa ctc c-3' (SEQ ID NO:20)
	g3883-1.6 rev	5'-tgt ttc aga gta gcc aat tc-3' (SEQ ID NO:21)
30	g13138-1.4 for	5'-cac gtt agt tag tta gaa gg-3' (SEQ ID NO:22)
	g13138-1.4 rev	5'-ctg atg ttc tct aca aat gg-3' (SEQ ID NO:23)
	B9-1.8 for	5'-cgt atc cgc att tct tca ctg c-3' (SEQ ID NO:24)
35	B9-1.8 rev	5'-cat ctg caa cat ctt ccc cag-3' (SEQ ID NO:25)
	IH1L-1.6 for	5'-ttg agt cct tct tgt ctg-3' (SEQ ID NO:26)

1H1L-1.6 rev	5'-cta gag ctt gaa agt tga tg-3' (SEQ ID NO:27)
5F7R-1.5 for	5'-gaa tgg tgt aac caa act c-3' (SEQ ID NO:28)
5F7R-1.5 rev	5'-cat acc gta tga tgg aac-3' (SEQ ID NO:29)
5	
20B4L-1.6 for	5'-gaa ctc att gta tgg acc-3' (SEQ ID NO:30)
20B4L-1.6 rev	5'-cta aga tgg gaa tgt tgg-3' (SEQ ID NO:31)
10	
8A6-1.3 for	5'-cca aga aga gaa aac gga ga-3' (SEQ ID NO:32)
8A6-1.3 rev	5'-aac aat agg agg tgc aga gt-3' (SEQ ID NO:33)

Primers to amplify across the *lsd1* deletion:

<i>lsd1</i> far side:	5'-acc taa caa aaa gaa aag tgt gtg agg-3' (SEQ ID NO:34)
<i>lsd1</i> outside	5'-ata ata aac cct act agc tct aac aag-3' (SEQ ID NO:35)
15 <i>lsd1</i> alt. spl. 5'	5'-ctg cta ctt tca tcc aaa c-3' (SEQ ID NO:36)

Example V Vector construction for complementation

The Agrobacterium vacuum infiltration procedure was used to generate transgenic plants (Bechtold et al., 1993; Grant et al., 1995). Vectors were derived from pGPTV-Hyg (Becker et al., 1992) as follows: pSGCGF was made by restricting pGPTV-Hyg with HindIII and SacI and replacing this fragment with a HindIII-SacI fragment containing the polylinker from pIC20H (GenBank accession L08912; provided by Steve Goff, Novartis, Research Triangle Park, N.C.). Either the 7kb XbaI or 4.5 kb PstI-XbaI genomic fragments were cloned into this, the former into the unique vector SalI site, the latter as a SacI-SalI fragment derived from an intermediate cloning step into pBS as a PstI-XbaI fragment. The pHyg35S vector was made by cloning a four enhancer-containing 35S promoter fragment as a HindIII-XbaI fragment into pGPTV-Hyg (provided by Dr. Douglas C. Boyes, Univ. of North Carolina, Chapel Hill). The EST 82D11 cDNA sequence was isolated as a SalI-XbaI fragment from pZL1 (Newman et al., 1994) and cloned into XbaI-XbaI digested pHyg35S.

Example VI Cloning

The genomic Ws-0 library in IGEM11 was a gift of Dr. Kenneth A. Feldmann (Univ. of Arizona). The cDNA library is an oligo-dT primed library prepared from polyA+ Col-0 mRNA from leaves cloned into LZAPII (Stratagene, La Jolla, CA) according to the manufacturer's instructions (gift of Dr. Douglas C. Boyes and Dr. Murray R. Grant).

Example VII LSD1 sequences

The sequences of the *LSD1* cDNA (SEQ ID NOS:14 and 15) and the 4.5 kb *LSD1* Xhol-PstI genomic fragment (SEQ ID NO:13; the longest 5'LSD1 cDNA starts at base 1892 of this sequence) are deposited in GenBank as accessions U 87833 and U 87834, respectively. Endpoints of the various *LSD1* cDNAs isolated are shown in Table 3A and examples are provided by SEQ ID NO: 14 (short form from cDNA MG7 as shown in Table 3) and SEQ ID NO:15 (long form, from cDNA MG8). The polypeptides deduced from these are shown in Fig. 11-12, respectively. Table 3B shows the sizes of each intron deduced from comparison of the sequence shown in SEQ ID NO:13.

10

Table 3. Sequence characteristics of the LSD1 gene
Endpoints of independent LSD1 cDNAs

<u>cDNA</u>	<u>5' end point</u>	<u>Alternate splice</u>	<u>3' end point</u>
MG7(2)	C 1	short	A 1021
EST 82D11	A 27	short	T 1031
MG4	C59	short	A 1188*
MG10	C 59	short	G 1225
MG5	G 67	short	A 1205
MG2 (4)	G 90	short	A 1106
MG8 (2)	G 98	long	A 1082
MG16 (2)	C 103	short	A 1066
MG11	C 117	long	G 1225

Numbers in parentheses refer to the number of isolates of the same clone. Nucleotide numbers at the 5' and 3' ends refer to nucleotide positions from SEQ ID NO:13. An A at the 3' endpoint can be either an A in the genomic sequence or the first A of the polyA tail. The endpoint marked with an * had no polyA tail.

Intron sizes

	<u>intron #</u>	<u>size in nucleotides</u>
30	1	88
	2 (short splice)	68
	2 (long splice)	129
	3	89
35	4	489
	5	100
	6	92

Intron splice junction positions are located at bses 198-199, 260-261, 447-448, 552-553, 692-693, 764-765, and 836-837 in SEQ ID NO:13.

5 Example VIII Genetic and physical mapping of *lsd1*

The *lsd1* mutation segregates as a monogenic recessive (Dietrich et al., 1994). F2 progeny of a cross between *lsd1* (Ws-0 background) and Col-0 (*LSD1*) were analyzed using the co-dominant amplified polymorphic sequences (CAPS) mapping procedure (Konieczny and Ausubel, 1993) to first establish linkage to the *AG* marker on chromosome 4. The closely linked g13838 probe (3 recombinants in 1632 meioses) was used to identify YAC (yeast artificial chromosome) clones (Schmidt et al., 1995; Schmidt et al., 1996). We constructed a physical contig of these YACs, shown in figure 1A. We used labeled YAC ends C1C1H1L, yUP5F7R and EG20B4L to isolate genomic phage clones, subcloned fragments from each of these, end-sequenced the subclones, derived primer sequences and developed new CAPS markers (see Tables 1 and 2). The CAPS markers 1H1L-1.6 and 5F7R-1.5 mapped closest to *lsd1* (1 and 3 recombinants, respectively from 2054 meioses); see Tables 1 and 2 for new CAPS markers. We hybridized these two CAPS markers to filters containing bacterial artificial chromosome (BAC) clone arrays (Choi et al., 1995, distributed by the Arabidopsis Biological Resource Center, Ohio St. Univ.), and isolated the five BAC clones depicted in Figure 2B. Because 5F7R-1.5 and 1H1L-1.6 genetically flank *lsd1* (Figure 1B), BAC clone 1G5 should contain the gene.

As 1G5 was the only BAC clone to physically span the relevant genetic region, we connected BACs 6H3 and 8A6 by walking in a genomic phage library. We defined a 5kb HindIII fragment from BAC 8A6 which hybridized only to itself and BAC 1G5. When used as a probe on filters containing restriction digests of the relevant BAC clones, this fragment hybridized to a 1.3 kb EcoRI fragment which also was present only on BACs 8A6 and 1G5. This 8A6-1.3 clone, (small box in Figure 1C) was used to isolate three phage clones, two of which are depicted in Figure 1C. Labeled inserts from each detected BAC clones 1G5, 6H3 and 8A6, thus providing multiple redundancy of genomic cloned DNA encompassing *lsd1*. We also converted 8A6-1.3 into a CAPS marker, and found that it cosegregated with *lsd1* in 2054 meioses. This map resolution of approximately 0.05 map units, suggested that *lsd1* was within 5-15 kb (at 100-300 kb per map unit; Schmidt et al., 1995; Schmidt et al., 1996) in either direction of 8A6-1.3.

We probed genomic Arabidopsis DNA blots of digested wild type Ws-0 and *lsd1* to confirm co-linearity of the cloned and genomic DNA immediately surrounding 8A6-1.3. We noted that a variety of fragments detected a genomic DNA rearrangement in *lsd1* relative to wild type Ws-0 (data not shown). This rearrangement corresponded to a loss of

restriction sites and a deletion as noted in Figures 1C and 3C. The *lsd1* mutant comes from an Agrobacterium mutagenized population of Arabidopsis, and it is known that the transformation procedure can generate non-T-DNA associated mutations (Feldmann, 1991). We subcloned and sequenced various wild type genomic DNA fragments at this position, and compared their sequences to several databases, including the Arabidopsis EST database (Rounsley et al., 1996, <http://www.tigr.org/tdb/at/at.html>). One EST clone (EST 82D11T7; GenBank accession T45220) exhibited blocks of identity to our genomic DNA sequence, suggesting the presence of introns in the latter. Because the gene encoding this EST is largely deleted in *lsd1*, it became a candidate *LSD1* gene.

10

Example IX Complementation of *lsd1*

15

To confirm that the genomic deletion encompasses *LSD1*, we constructed subclones from the genomic phage as shown in Figure 3C for complementation into the T-DNA binary vector pSGCGF. Because the typical method for generation of transgenic Arabidopsis, vacuum infiltration of Agrobacterium carrying binary T-DNA vectors, triggers the propagative cell death indicative of the *lsd1* phenotype, we devised an alternate complementation strategy. We transformed F1 plants of *lsd1* x Col-0, and plated surface-sterilized seeds of the next (F2) generation onto media containing hygromycin as a selective antibiotic. We then identified hygromycin resistant transformants which were homozygous for Ws-0 alleles at 5F7R-1.5, 1H1L-1.6, and 8A6-1.3, and thus were *lsd1/lsd1* homozygous mutants. These individuals contained both mutant and wild type alleles for the CAPS marker which spans the *lsd1* deletion, because a wild type allele is present on the transgene. These transgenic plants were treated with droplets of 2,6-dichloroisonicotinic acid (INA); 0.3 mg/ml wettable powder containing 25% active ingredient, Uknas et al., 1993a) a potent inducer of SAR and the *lsd1* phenotype (Dietrich et al., 1994). If the mutation were complemented, then INA treatment should not lead to spreading cell death. Table 4 shows that transgenic plants carrying either the 7kb XhoI fragment or the 4.5 kb PstI-XhoI (Figure 3C) all survived this treatment, and are thus complemented for the *lsd1* mutation. Selfed F3 progeny from a complemented F2 individual carrying either the 4.5 kb XhoI-PstI fragment or the 7 kb XhoI fragment were also analyzed. All F3 progeny which inherited the transgene were complemented (Table 4), while all of their non-transgenic sibs still exhibited the *lsd1* phenotype (data not shown). In no case did wild type control plants exhibit spreading cell death after INA application.

35

Table 4. Complementation of the *lsd1* mutant

of plants complemented/# transgenics tested from:

<u>Construct</u>	<u>Independent F2s</u>	<u>Transgenic F3 progeny</u>
------------------	------------------------	------------------------------

7 kb Xhol	1/1 ^A 3/3 ^C	20/20 ^B 21/21 ^C
kb PstI-Xhol	2/2 ^A	14/14 ^B
35S-cDNA	1/1 ^A	19/19 ^B

5 ^A Selected for hygromycin resistance and screened for homozygous Ws-0 alleles through the *lsd1* genetic interval as described, except where noted in ^C. Individual F₂s were both drop tested with INA and shifted to LD conditions (Dietrich et al., 1994).

10 ^B Selfed progeny from a complemented F₂ individual (homozygous Ws-0 alleles through the *lsd1* interval) were screened by PCR at F₃ for presence of the hygromycin resistance gene and then INA tested.

15 ^C F₂ parents were identified as hygromycin resistant and heterozygous through the *lsd1* interval, then selfed and re-screened as hygromycin resistant and homozygous Ws-0 through the *lsd1* interval at F₃ before INA testing.

20 Due to low numbers of independent F2 transformants which were homozygous mutant through the *lsd1* interval from the original transformation, we also isolated F2 transformants carrying the 7 kb XhoI fragment which were originally identified as heterozygote at the CAPS markers flanking *lsd1*. Selfed progeny from these should segregate both the transgene and the *lsd1* mutation. Among these progeny, we identified F3 individuals which were homozygous Ws-0 through the *lsd1* interval and carried the transgene. As shown in Table 4, these also were all complemented for protection against INA-induced spreading cell death. We conclude that the 4.5kb PstI-XhoI fragment carries the *lsd1* gene and sufficient *cis* control elements to ensure its expression.

25 All transgenic plants complemented for the INA-induced *lsd1* mutant phenotype were also complemented for initiation of spreading cell death after transfer to non-permissive long day conditions as well (Dietrich et al., 1994; not shown). Thus, the complementing DNA corrects the mutant phenotype induced by two independent stimuli.

Example X Identification of alternately spliced LSD1 transcripts

30 We sequenced all of the complementing 4.5 kb PstI-XhoI genomic DNA fragment (SEQ ID NO:13), eight independent cDNAs (Example VII) and completed the sequence of the full 82D11T7 EST sequence. Among the cDNAs, we identified two classes expressing open reading frames of either 184 or 189 amino acids (SEQ ID NO:16 and 17). An alternate splice which adds 61bp to the 5' region of some cDNAs also provides an alternate translation start, hence, the extra five amino acids in SEQ ID NO:17. The sequences of both cDNA classes matched exactly the genomic sequence except at the positions of 7 introns

(see Table 3). Nucleotide 1 of the longest cDNA is at position 1892 in the 4.5 kb PstI-XbaI genomic sequence (SEQ ID NO:13). Thus, 1891 nucleotides of promoter are sufficient for appropriate expression in complementation of the *lsd1* mutation. The cDNA 5' ends are clustered (Table 3), suggesting that the longest could be full length. We also complemented the *lsd1* mutation by transformation of the full insert from EST clone 82D11T7 expressed from the strong and constitutive cauliflower mosaic virus 35S promoter (see Table 3) proving that this cDNA contains the entire LSD1 coding region. The 3' ends of these cDNAs are very heterogeneous, suggesting the presence of multiple polyadenylation addition signals (Table 3). No other significant open reading frames were observed in the 4.5 kb PstI-XbaI genomic clone.

When either the EST 82D11T7 clone, or a 0.8 kb EcoRI-XbaI genomic fragment covering the *lsd1* deletion were used as probes on RNA blots, a rare mRNA of approximately 1.2 kb was detected in leaf tissue of wild type Ws-0 plants (Figure 3). This length is consistent with the size of the longest cDNA, supporting the conclusion that we have identified a nearly full-length transcript. Importantly, this mRNA was completely lacking in mRNA prepared from *lsd1* leaves, furthering the argument that it encodes LSD1. The finding that *lsd1* is an mRNA allele was corroborated by sequencing across the genomic deletion in the mutant (Figure 3). The 5' border of the deletion is an A at nucleotide 55 and the 3' boundary is in the fourth intron (data not shown). It is noteworthy that expression of this candidate mRNA was unaffected by application of INA (Figure 4, top). The expected high level of INA-induced PR-1 mRNA accumulation in leaves of both wild type and *lsd1* (Figure 4, middle) served as a control in this experiment for efficacy of INA treatment.

The *lsd1* phenotype can be observed in all cell types examined after initiation of lesion formation (Dietrich et al., 1994). RNA blot analysis of seedlings, stems, leaves and flowers demonstrated that the *LSD1* gene is expressed constitutively in each of these *Arabidopsis* tissues (data not shown). Thus, the requirement for *LSD1* activity in these tissues is consistent with the gene's expression pattern.

30 Example XI The *LSD1* mRNA encodes a novel zinc-finger domain

We searched a variety of databases with the predicted translation product of the *LSD1* cDNA sequence. Several striking features emerged. First, there are three zinc-finger domains, depicted in Figure 5 (SEQ ID NOS:1-3), which share remarkable homology with one another. These are C-x-x-C, or type IV, zinc-fingers, according to the classification of Sánchez-García and Rabbitts (1994), and they share most homology with plant relatives of the GATA-1 transcription factor (Evans and Felsenfeld, 1989; Omichinski et al., 1993). The plant members of this sub-family described to date include the *CO* gene, which controls

transition to flowering (Putterill et al., 1995), a set of related DNA binding proteins (Yanagisawa, 1995; De Paolis et al., 1996) and a gene whose transcription is salt stress-induced (Lippuner et al., 1996). None of these proteins shares with LSD1 the consensus homology within the Zn-fingers. The second homology domain is derived from the carboxyl portion of LSD1, from residues 129 to 180 (Figure 6-SEQ ID NO:4). This region of LSD1 exhibits homology to three broad classes of regulatory proteins. First, all mammalian insulin receptor substrates; second, a set of animal transcription factors; and third, a maize transcription initiator binding protein.

The conceptual LSD1 translation product also identified two additional *Arabidopsis* ESTs via their predicted amino acid homology. Importantly, each has at least one C-x-x-C Zn-finger and most of the associated consensus residues found in the LSD1 internal homologies. They are ESTs 172A7T7 (GenBank R6552)(SEQ ID NO: 58 and 132J21T7 (GenBank T45809). Thus, it is probable that LSD1 is the first member of a widely distributed Zn-finger sub-family in plants, defined by the internal homology within each zinc-finger. The other amino acids in the consensus section are not known to be found in any other zinc finger proteins.

Example XII Identification of expressed target sequence tags (EST) and cDNAs containing LSD1-type zinc finger domains

As discussed in the text prior to the Examples, the predicted amino acid sequence of the LSD1 zinc fingers was used to search the GenBank database (NCBI). Two *Arabidopsis thaliana* ESTs (EST132J21T7 and EST 172A7T7) were identified, each of which contains at least two zinc finger domains and most of the associated consensus residues found in the LSD1 internal homologies (Dietrich, 1997). These ESTs were ordered from Ohio State University Arabidopsis Biological Resource Stock Center and resequenced. Sequences were analyzed with the Genetics Computer Group programs (Devereaux et al., 1994). A specific probe isolated from EST172A7T7 was subsequently used for screening of cDNA and genomic libraries. The bacterial strain carrying EST132J21T7, however, was not viable. Therefore, degenerated primers were designed based on the EST132J21T7 sequence. Genomic *Arabidopsis thaliana* Ws-0 DNA was used in the PCR reaction and gave rise to a specific PCR product of approximately 400 bp. This fragment was subcloned via the TA Cloning Kit (Invitrogen, Carlsbad, CA) into pBluescript KS(+). Two new genes were identified as described here. Their predicted protein products are highly related to that of LSD1 indicating an involvement in the control of cell death in plants

35

Example XIII LOLI cDNA

Poly A + RNA isolated from uninduced and *P. syringae* DC3000 induced

Arabidopsis thaliana Col-0 leaf tissue was reverse transcribed. The resulting cDNA population was subcloned unidirectionally into the EcoRI/Xhol - sites of a lambda-Zap II vector using the cDNA-synthesis Kit (Stratagene, La Jolla, CA) according to the manufacturer's directions. The titer of this MG-library was calculated as 2.5×10^6 pfu

5 Approximately 8×10^5 pfu of the amplified MG-library were subsequently screened with α^{32} P dCTP labeled probes (Stratagene 'Prime it' Kit) specific for EST132J21T7 or EST172A7T7. With the probe specific for EST132J21T7, four cDNA clones were identified and subcloned via the Stratagene excision system. One clone contained an insert of less than 100 bp in length and was not further analyzed. The three remaining clones were

10 sequenced by standard protocol (primers: M13F, M13R, PE6, and PE7; for primer sequences refer to Table 5, below). Clones 2 and 3 contained identical open reading frames (ORFs) and were homologous to EST132J21T7 and to another identical and overlapping EST clone, EST119C9T7. The fourth clone consisted of a chimeric cDNA of approximately 1500 bp, with approximately 400 bp similarity to EST132J21T7,

15 EST119C9T7, and clones 2 and 3. It was also not analyzed further.

Table 5. Primers and primer sequences used

<u>Primer</u>	<u>Primer Sequence</u>	<u>SEQ ID NO:</u>
M13F	5'- GTA AAA CGA CGG CCA TG -3'	37
M13R	5'- GGA AAC AGC TAT GAC CAT G -3'	38
PE6	5'- TTC ATG GCA ATG GTG TGA CCC C -3'	39
PE7	5'- CTG CCG GAT TCT TGA TCG AAG A -3'	40
PE8	5'- AGA GGA AGG TCC GCC TCC GG -3'	41
25 PE9	5'- CTC TGC TCT CCT GAG ACT GCT T -3'	42
PE13	5'- CAT CAT AAT GTC TCC TTT TGA GAC -3'	43
PE15	5'- GCC ATC CAT TAT TCA TCG CCT -3'	44
PE23	5'- GAG GAG GAA GAA CTG CAG ATT CC -3'	45
PE30	5'- GTG CTC CAT GTC CAA ATC ATA C -3'	46

30

Clone 2, with an insert length of 908 bp represents a full length cDNA clone, as determined by the presence of an open reading frame flanked by untranslated sequences, and was renamed *LOL1* (*Lsd one like*)(SEQ ID NO:47). We confirmed that the *LOL1* cDNA and EST132J21T7 are encoded by the same gene using genomic DNA (Southern) blot analysis (data not shown). The LOL1 protein of 154 amino acids (SEQ ID NO:48) contains three zinc finger domains of the LSD1-type (SEQ ID NOS:49-51). The consensus

sequence of the LOL1 zinc finger domains is defined by CxxCxxLLMYxxGAXsxCxxC (SEQ ID NO:53).

Example XIV LOL2 cDNA

5 By screening the MG-cDNA-library, no clones homologous to EST172A7T7 could be obtained. Therefore, the AB-cDNA-library (derived from RNA isolated from different tissues of sterile grown plants, available at the European Arabidopsis Stock Center, Cologne, Germany) was screened with α^{32} P dCTP labeled probe specific for EST172A7T7. Six homologous cDNA clones were obtained and subcloned into the SmaI site of
10 pBluescript KS(+). Restriction analysis indicated that the inserts were encoded by the same gene. Only the longest insert was sequenced following standard protocol (primers used: M13F, M13R, PE8 and PE9: for primer sequences, refer to Table 5. We demonstrated that this insert contained an ORF of 500 bp homologous to EST172A7T7. This non-full length cDNA was designated *LOL2* (SEQ ID NO:54). The deduced protein (SEQ ID NO:55)
15 consisting of two LSD1-type zinc finger domains extending from bases 130-195 and 244-309 of SEQ ID NO:54 (SEQ ID NOS:56-57, respectively). Comparison to EST172A7T7 shows that the EST (SEQ ID NO:58) contains a 124 bp insertion (bases 386-509 after the second zinc finger of SEQ ID NO:58), leading to a different C-terminal. Comparison of these two partial cDNA sequences with the genomic *LOL2* sequence (see below)
20 demonstrates that they are alternate splice forms from the same gene encoding two related proteins. This conclusion is strengthened by the fact that the *LOL2* cDNA and EST172A7T7 hybridize to the same genomic DNA fragment and therefore are encoded by the same gene (data not shown). Thus, sequence analysis of genomic *LOL2* clones shows that the non-identical C-termini of *LOL2* and EST172A7T7 are due to alternative splice
25 sites. The genomic sequence of *LOL2* (SEQ ID NO:59, has a putative TATA-box sequence and polyadenylation signal (bases 922-930 and 2539-2544), and the exon borders of an alternative splice site (bases 2256-2382). The derived amino acid sequence extends from bases 1231-2462.

30 **Example XV Isolation of genomic *LOL2* sequences from an *Arabidopsis thaliana* Col-0 library**

35 8×10^5 genomic lambda clones (lambda GEM11, European Arabidopsis Stock Center) were screen with a α^{32} P dCTP labeled probe specific for EST172A7T7. Nine clones homologous to *LOL2* EST172A7T7 could be identified. Restriction analysis demonstrated that the nine clones belonged to five different classes. Inserts ranging from two to five kb in size were isolated and subcloned into either SacI or BamHI sites of pBluescript KS(+). Sequence information derives from two overlapping clones,

sequentially sequenced with primers M13R, PE9, PE13, PE15, PE23 and PE30 (see Table 5).

The genomic *LOL2* sequence has a length of 3060 bp. Promoter and 5' untranslated regions consist of approximately 1200 bp. The translation products are encoded by three 5 exons, which are interrupted by two introns of 182 bp and 458 bp length, respectively. The overall length of the coding sequence is 1232 bp. Due to alternative splice sites, two proteins which differ in their C-terminal regions are encoded by the *LOL2* gene (SEQ ID NO:59). A first protein, of 155 amino acids (SEQ ID NO:60), is identical to the *LOL2* cDNA and contains two zinc finger domains of the LSD1-type. The other translation 10 product corresponds to EST172A7T7, consists of 147 amino acids, and contains two and a half zinc finger domains (SEQ ID NO:61). The consensus sequence of the two zinc finger domains of LOL2 is CxxCxxLLxYxxGxxxVxCSSC (SEQ ID NO:62).

Example XVI Obtaining interacting genes

15 The methodology for this Example is known to those skilled in the art and summarized in Fields and Sternglanz, 1994, and Finley and Brent, 1996. The *LSD1* short or *LDS1* long open reading frames were cloned into the "bait vector" pEG202 of the commonly available LexA yeast two-hybrid system (MatchmakerTM, Clonetech, Palo Alto, CA) to generate plasmids pEG202-L and pEG202-S. These encode fusion proteins of the LexA DNA binding domain and the full length LSD1 protein of both long and short isoforms (SEQ ID NOS 14 and 15). Yeast strain EGY48 is transformed with this plasmid, and appropriate controls performed to ascertain the LSD1 fusion protein encoded by 20 plasmids pEG202-L and pEG202-S did not intrinsically activate expression of the yeast markers used in this system. A yeast gene expression library was constructed in plasmid pJG4-5 using RNA from *Arabidopsis* leaves infected with *Pseudomonas syringae*. This library encodes fusion proteins of expressed *Arabidopsis* genes and the B42 transcriptional activation domain. The library was transformed en masse into the yeast strain EGY48 25 carrying either plasmid pEG202-L or -S. From an equivalent of 6 million clones screened, 122 were isolated. The longest insert of a member from each of these classes was sequenced using standard DNA sequencing methods. Because the novel *Arabidopsis* gene so identified is produced as an active translation fusion in this system, one is immediately able to identify the deduced protein sequence. The most interesting sequences thus defined, 30 and their deduced protein sequences, are set forth herein as SEQ ID NOS: 66-91.

35 The first main class of LSD1-interacting proteins has no database homologues. These proteins encode putative "sequestration" proteins for LSD1 whose function is to inhibit LSD1 function until the correct pathogen signal is received. Their utility lies in

manipulation of the interaction with LSD1 in plant cells such that LSD1 is altered in its ability to regulate the response to pathogen. Alternatively, these novel LSD1-interacting proteins may encode new components of the gene regulation machinery working together with LSD1 to control transcription in response to pathogen infection. These proteins are 5 valuable because of the knowledge that LSD1 is a key regulator of cell death in plants in response to pathogens. Proteins which physically interact with LSD1 share in this cellular function.

The second class defines proteins having database homologies to other proteins, strongly suggesting a role in control of gene transcription (e.g., CAAT box binding proteins 10 which are known to bind the common CAAT regulatory unit in DNA preceding nearly all genes encoding eukaryotic mRNA). This finding is completely consistent with the embodiment described above, in which the LSD1 partner proteins identify other components of the gene regulatory machinery required for response to pathogens. Manipulation of the expression of, for example, CAAT box binding proteins, will result in 15 altered response to pathogen infection.

While the invention has been described with reference to specific embodiments, it will be appreciated that numerous variations, modifications, and embodiments are possible, and accordingly, all such variations, modifications, and embodiments are to be regarded as 20 being within the spirit and scope of the invention.

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SEQUENCE LISTING**SEQ ID NO:1**

LVCHGCRNLLMYPRGASNVRCALCNTINMV

SEQ ID NO:2

IICGGCRTMLMYTRGASSVRCSCCQTTNLV

SEQ ID NO:3

INCGHCRTTLMYPYGASSVKCAVCQFVTNV

SEQ ID NO:4

MSNGRV-PLPTNRP-NGTACPPST-STSTPPSQQTQTVVVENPMSVDESGKLVSNV

SEQ ID NO:5

MSPG-VAPVPSNRKGNGDYMPMSPKSVSAP-QQIINPIRRHPQRVDPNGYMM

SEQ ID NO:6

VPLPTNRP-NGTACPPSTSTSTPPSQQTQTVVVENPMSVDESGKLVSNV

SEQ ID NO:7

VPLPANNPVV-TTVVPSTPPSQPPAVCPPVV

SEQ ID NO:8

VPLPANNPVV-TTVVPSTPPSQPPAVCPPVV

SEQ ID NO:9

IPVYTNSNV-GTALPPSVSPSVSPSVT

SEQ ID NO:10

VVLP-NAAPAGAAAPPSSRSTS

SEQ ID NO:11

SNGRVPLPTNRPN-GTACPPSTSTSTPPSQQTQTVVVENPMSVDESGKLVSNV

SEQ ID NO:12

SRALVPVPAADPNAG-AIVPANKSKRSPEQQRRIRR

SEQ ID NO:13

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 730 750 770
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 970 990 1010
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 2290 2310 2330
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 2350 2370 2390
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 2590 2610 2630
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 2650 2670 2690
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 2710 2730 2750
 TATCGTGACATCCATATCAATCCTTTAAAGACCATGTATTATATTGCTTTATAAGGTCT

 2770 2790 2810
 TTAGTCCTTAAAGAACATTCTTCACACTTTGTTGATAACATTGTTCTGTGGAGATGA

 2830 2850 2870
 TGCTTACGTAACGTATTCACCTTCCCAGATGTATATGAATTCTGAATTCTGAAAAT

 2890 2910 2930
 ATCTGGGATTTGTAAAGCAGCTGAAAGTACTAAACAAAGCTTTAGATGGTCCCAGGTG

 2950 2970 2990
 GACTAGGTAACTACTTGTAGAGCTAGTAGGGTTATTATGTTGTTGATCTACCAT

 3010 3030 3050
 TAGATTCTTATCTTAATTAGCGTCTAACGCTGTCATTAGCTGTATGATTATCATT

 3070 3090 3110
 ATCCATGACTGCTTAAGAACATTGCTGATTACCTCGTCATTAGTATTCTGGATTTT

 3130 3150 3170
 CTAGCATTAACATTGCTTGTCTGAATCTGTGCGTGTCTTTTGAAATCGACAGCGC

 3190 3210 3230
 ACTCCAATCAGGTTGCCATGCTCCTCCAGTCAGGTTGCGCAAATCAATTGTGGGCATT

3250 3270 3290
 GTCGGACGACCCTCATGTATCCTTACGGTCATCATCCGTCAAATGCGCTGTTGTCAAT

 3310 3330 3350
 TCGTAACCTAACGTTAATGTGATTATTCCCTATCTATTAAAGCCACCTCTGCATGGTTGAGTT

 3370 3390 3410
 AAGTATAGAGATCTTCTGTTGAAATTTCATTTCTGATTCAATTGCACTCCTTAGATG

 3430 3450 3470
 AGCAATGGAAGGGGTACCTCTCCCAACTAACCGGCCAATGGAACAGCTTGTCCCCCTC

 3490 3510 3530
 TACATCAACTGTGAGTTATCAAATTATGAATTGTAATAGTTCTGTATATTCTTATGGAA

 3550 3570 3590
 CTGGTACTTACTCTGTTCATCGATTTTACCAACAGTCAACACCACCCCTCAG

 3610 3630 3650
 ACCCAAACCGTTGTTGAGAAAACCCATGTCGTTGATGAAAGCGGAAAGTTGGTGAGT

 3670 3690 3710
 ATTTCTATCACCTGTGTTCTCTTCTTATTACACATTAGAGGAAGATATGACAAAGTG

 3730 3750 3770
 ACTGAAACACACAATTGCAGGTGAGCAATGTTGTTGGAGTGACAACGTGACAAAAAG

 3790 3810 3830
 TAATCAAGAATGAGTGAGATCTAAAGATCAAATCCAATTCTCCTCTATTCCCTGCGTT

 3850 3870 3890
 TGGTTTGTGCATATTACATACGCGGAAAACGTATGTTATATCTCTTGACTCCTTTT

 3910 3930 3950
 TAACCCAAGAGAAAAAGCTTATCAGAATCTCTGTTACTGCATTATTGGGGTTATTCAA

 3970 3990 4010
 AGTTGAAGACACAAAGTTTGTCTGAATAATTGGCATTCTTGCTCCATGGAACCTTG

 4030 4050 4070
 ACCTTCTCTCTGTTAGTTGACTCTAAACTCCATGGCCCTGTGGCATTGTTAATGT

 4090 4110 4130
 ATGTATGAATATAATCTGATAACACCAACCAATCATTAAGATTGGGTTGAAATCTGTCT

 4150 4170 4190
 CTTCCGTGGATGAGATATGCTACATGTCACAAGAACTGGTCTTAGCTTGGTAGATAAGA

 4210 4230 4250
 CTTGTCTTAGAAGCAAGTCTTGAATCTGAAATCTGGAAATCTATTGCAAGTAATCTGTCACAAC

 4270 4290 4310
 AACCATAACCTAATCAGTCAGTACCCCTCCAAGAAACATTAAGTTAGATGATCCGACAAA

 4330 4350 4370

ACCTCTCAACAAAACCAACTCTTCCATATAAATCTTTAACACTGGACCAAATTNC
 4390 4410 4430
 ACCCTTCCTCTTGATCCTCCCTGCATCACAAATGGCCAAAAAAATGGTGGTGGCNGG
 4450 4470 4490
 TGGGTACCACAAAGAGCTGGAAACTACTCTTGGGCTGAGAATATTCATTGCATTGCTA
 4510
 CTTTAGCTGCAG

SEQ ID NO:14

10 30 50
 CTTACCGCGTCATGTAAAAAAAAAGAAGCGTAAATTACGAAAAACAGAGAGATAATCCG
 70 90 110
 GGCATTGAGATTTGGAGATAGAGAGAGAGAGAAAAATCGAAATCTATTGTCTATCTCCTCA
 130 150 170
 ATTTGGATTGGATTTCTGCATATCATCGCTCTAGATTCGCGGGTTTGGATTGCGATT
 190 210 230
 CTTACCCCTCTCCAATCGAAGTTTGGCTTGAATTGGATTGGTTTCGTTCCAAAAT
 250 270 290
 CAGCTCTTTGTTAATCAGATATGCAGGACCAGCTGGTGTGTAGGGTTGAGAATTT
 310 330 350
 ATTGATGTATCCTAGAGGAGCATCTAATGTGCGTTGCGTTATGTAACACTATCAACAT
 370 390 410
 GGTTCCCTCCTCCCTCCACCTCACGACATGGCACACATTATATGTGGTGGTTGTAGAAC
 430 450 470
 GATGCTTATGTATAACGCGTGGGGCTAGTAGCGTAAGATGTTCTGCTGTCAAACACTACGAA
 490 510 530
 CCTTGTGCCAGCGCACTCCAATCAGGTTGCCATGCTCCCTCAGTCAGGTTGCGCAGAT
 550 570 590
 CAATTGTGGGCATTGCGGACGACCCCTCATGTATCCTTACGGTGCATCATCCGTCAAATG
 610 630 650
 CGCTGTTGTCATTGTAACGTTAATATGAGCAATGGAAGGGTACCTCTCCCAAC
 670 690 710
 TAACCGGCCAAATGGAACAGCTTGTCCCCCTCTACATCAACTTCAACACCCACCTCTCA
 730 750 770
 GACCCAAACCGTTGTTGAGAAAACCCATGTCGTTGATGAAAGCGGAAAGTTGGTGAG
 790 810 830
 CAATGTTGTTGGAGTGACAACGTGACAAAAAGTAATCAAGAATGAGTGGAGATCTTAAA

850 870 890
 GATCAAATCCAAATTCTTCCTCTGTTCTGCCTGGTTGTGCATATTACATACGCGGA
 910 930 950
 AAAACTGTATGTTATATATCTCTTGAUTCCTTTAACCCAAGAGAAAAAGCTTATCAGA
 970
AAAAAAAAAAAAAAA

SEQ ID NO:15

10 30 50
 GAAATCTATTGTCTATCTCCTCAATTGGATTGGATTTCTGCATATCATCGCTCTAGCT
 70 90 110
 TTTCGCGGGTTTGGATTGATTCTACCCCTCTCCAATCGAAGTTTGGCTTGATT
 130 150 170
 GGATTTGGGTTTCGTTCCAAAATCAGCTTTTGTAAATCAGGGTTCATCTGTGTGGG
 190 210 230
 TCTTGTTTGAAGCAATTGTGTGTTGGATGAAAGTAGCAGATATGCAGGACCAGCT
 250 270 290
 GGTGTGTATGGTTGTAGGAATTATTGATGTATCCTAGAGGAGCATCTAATGTGCGTTG
 310 330 350
 TGCGTTATGTAACACTATCAACATGGTCCTCCTCCACCTCACGACATGGCACA
 370 390 410
 CATTATATGTGGTGGTTGTAGAACAAATGTTATGTATAACGCGTGGGGCTAGTAGCGTAAG
 430 450 470
 ATGCTCTGCTGTCAAACACTACGAACCTTGTGCCAGCGCACTCCAATCAGGTTGCCATGC
 490 510 530
 TCCCTCCAGTCAGGTTGCGCAGATCAATTGTGGGCATTGTCGGACGACCCCTCATGTATCC
 550 570 590
 TTACGGTGCATCATCCGTCAAATGCGCTGTTGTCAATTGTAACGTTAACGTTAATATGAG
 610 630 650
 CAATGGAAGGGTACCTCTCCAACTAACCGGCCAATGGAACAGCTTGTCCCCCTCTAC
 670 690 710
 ATCAACTCAACACCACCCCTCTCAGACCCAAACCGTTGTAGAAAACCCCAGTCCGT
 730 750 770
 TGATGAAAGCGGAAAGTTGGTGAGCAATGTTGTTGGAGTGACAACGTGACAAAAAGTA
 790 810 830
 ATCAAGAATGAGTGAGATCTTAAAGATCAAATCCAAATTCTCCTCTATTGCGTTG
 850 870 890
 GTTTGTGCATATTACATACGCGGAAAAACTGTATGTTATATCTCTTGAUTCCTTTTA

910	930	950
ACCCAAGAGAAAAAGCTTATCAGAATCTCTGTTACTGCATTATTGGGGTTTATTCAAAG		
970	990	
TTGAAGACACAAGGTTTGCTCGAAAAAAAAAAAAAA		

SEQ ID NO:16

MetGlnAspGlnLeuValCysHisGlyCysArgAsnLeuLeuMetTyrProArgGlyAla	10	20
SerAsnValArgCysAlaLeuCysAsnThrIleAsnMetValProProProProPro	30	40
HisAspMetAlaHisIleIleCysGlyGlyCysArgThrMetLeuMetTyrThrArgGly	50	60
AlaSerSerValArgCysSerCysCysGlnThrThrAsnLeuValProAlaHisSerAsn	70	80
GlnValAlaHisAlaProSerSerGlnValAlaGlnIleAsnCysGlyHisCysArgThr	90	100
ThrLeuMetTyrProTyrGlyAlaSerSerValLysCysAlaValCysGlnPheValThr	110	120
AsnValAsnMetSerAsnGlyArgValProLeuProThrAsnArgProAsnGlyThrAla	130	140
CysProProSerThrSerThrProProSerGlnThrGlnThrValValValGlu	150	160
AsnProMetSerValAspGluSerGlyLysLeuValSerAsnValValValGlyValThr	170	180
ThrAspLysLys		

SEQ ID NO:17

MetLysValAlaAspMetGlnAspGlnLeuValCysHisGlyCysArgAsnLeuLeuMet	10	20
TyrProArgGlyAlaSerAsnValArgCysAlaLeuCysAsnThrIleAsnMetValPro	30	40
ProProProProProHisAspMetAlaHisIleIleCysGlyGlyCysArgThrMetLeu	50	60
MetTyrThrArgGlyAlaSerSerValArgCysSerCysCysGlnThrThrAsnLeuVal		

70

80

ProAlaHisSerAsnGlnValAlaHisAlaProSerSerGlnValAlaGlnIleAsnCys
90 100

GlyHisCysArgThrThrLeuMetTyrProTyrGlyAlaSerSerValLysCysAlaVal
110 120

CysGlnPheValThrAsnValAsnMetSerAsnGlyArgValProLeuProThrAsnArg
130 140

ProAsnGlyThrAlaCysProProSerThrSerThrSerThrProProSerGlnThrGln
150 160

ThrValValValGluAsnProMetSerValAspGluSerGlyLysLeuValSerAsnVal
170 180

ValValGlyValThrThrAspLysLys

SEQ ID NO:18

5'-CAG TGG ATC TTT CCT CAG ACG-3'

SEQ ID NO:19

5'-CAT CTT CTT CTG CAA TCT GGG-3'

SEQ ID NO:20

5'-CAT CCA TCA AAC AAA CTC C-3'

SEQ ID NO:21

5'-TGT TTC AGA GTA GCC AAT TC-3'

SEQ ID NO:22

5'-CAC GTT AGT TAG TTA GAA GG-3'

SEQ ID NO:23

5'-CTG ATG TTC TCT ACA AAT GG-3'

SEQ ID NO:24

5'-CGT ATC CGC ATT TCT TCA CTG C-3'

SEQ ID NO:25

5'-CAT CTG CAA CAT CTT CCC CAG-3'

SEQ ID NO:26

5'-TTG AGT CCT TCT TGT CTG-3'

SEQ ID NO:27

5'-CTA GAG CTT GAA AGT TGA TG-3'

SEQ ID NO:28

5'-GAA TGG TGT AAC CAA ACT C-3'

SEQ ID NO:29

5'-CAT ACC GTA TGA TGG AAC-3'

SEQ ID NO:30

5'-GAA CTC ATT GTA TGG ACC-3'

SEQ ID NO:31

5'-CTA AGA TGG GAA TGT TGG-3'

SEQ ID NO:32

5'-CCA AGA AGA GAA AAC GGA GA-3'

SEQ ID NO:33

5'-AAC AAT AGG AGG TGC AGA GT-3'

SEQ ID NO:34

5'-ACC TAA CAA AAA GAA AAG TGT GTG AGG-3'

SEQ ID NO:35

5'-ATA ATA AAC CCT ACT AGC TCT AAC AAG-3'

SEQ ID NO:36

5'-CTG CTA CTT TCA TCC AAA C-3'

SEQ ID NO:37

5'- GTA AAA CGA CGG CCA TG -3'

SEQ ID NO:38

5'- GGA AAC AGC TAT GAC CAT G -3'

SEQ ID NO:39

5'- TTC ATG GCA ATG GTG TGA CCC C -3'

SEQ ID NO:40

5'- CTG CCG GAT TCT TGA TCG AAG A -3'

SEQ ID NO:41

5'- AGA GGA AGG TCC GCC TCC GG -3'

SEQ ID NO:42

5'- CTC TGC TCT CCT GAG ACT GCT T -3'

SEQ ID NO:43

5'- CAT CAT AAT GTC TCC TTT TGA GAC -3'

SEQ ID NO:44

5'- GCC ATC CAT TAT TCA TCG CCT -3'

SEQ ID NO:45

5'- GAG GAG GAA GAA CTG CAG ATT CC -3'

SEQ ID NO:46

5'- GTG CTC CAT GTC CAA ATC ATA C -3'

SEQ ID NO:47

10

30

50

AATATATCGAAACGAGATTCCACAATTAGTCTCTAGTCAGAGCTTCATGGCAATGGTG

70

90

110

TGACCCCAAATATAGATTGATGAAAGTGAGGAAATAGGAGAAGAAATGAAGAACACAGG
 130 150 170
 ATGTGTCTTCTTCTAAGTCACTAACAAAATCAACAAAGAGGAGAACGCCATTATTATA
 190 210 230
 TAATAGAGAGATTGAGAGAAGAGATTATCaaaaaaaaATTGCAATTCTTCTGGAGTG
 250 270 290
 ATAATGCCAGTCCCTCTGCACCATATCCAACACCTCCGGCACCGGCACAGGCTCCGTC
 310 330 350
 GTACAAACACTCCTCCGGCAAATGGAAGTACAAGTGGCAGAGCCAGTTAGTGTGTTCAAGG
 370 390 410
 TTGCAGAAACCTCTGATGTATCCCCTCGGAGCAACCTCCGTCTGCTGCGCCGTCTGTAA
 430 450 470
 CGCCCGTCACGGCCGTTCCCTCCGCCGGAACGGAGATGGCACAGTTAGTATGTGGAGGATG
 490 510 530
 CCATACACTCTTAATGTACATTCTGGAGCTACAAGTGTCAATGTTCTGTTGTACAC
 550 570 590
 TGTTAATCTGCCCTCGAACGCAACCAAGTAGCGCATGTGAATTGCGGAAACTGCATGAT
 610 630 650
 GCTACTAATGTATCAATATGGAGCAAGATCAGTGAAATGTGCCGTTGTAACCTTGTAC
 670 690 710
 ATCTGTTGGGGTTCAACGAGCACGACTGATTGAAAGTTAACATTAAACTTGGATCT
 730 750 770
 ATCTACCTATCAATATCTATTGAGTTATGAGCAATATAGAGGAAGCATCAAATCTTTTC
 790 810 830
 ACTCTCTTCGATCAAGAACCGCAGTTATGAGTTGAAACCATTTCGGAAGTAAAT
 850 870 890
 GAAATATGTAATTGTCGAAATTCTGACTTTGGTCTCTTGTCCGTTGATAGAGCTA
 910
 AAAAAAAA

SEQ ID NO: 48

Met Pro Val Pro Leu Ala Pro Tyr Pro Thr Pro Pro Ala Pro Ala Gln Ala Pro Ser Tyr
 10 20

Asn Thr Pro Pro Ala Asn Gly Ser Thr Ser Gly Gln Ser Gln Leu Val Cys Ser Gly Cys

30

40

ArgAsnLeuLeuMetTyrProValGlyAlaThrSerValCysCysAlaValCysAsnAla
 50 60

ValThrAlaValProProProGlyThrGluMetAlaGlnLeuValCysGlyGlyCysHis
 70 80

ThrLeuLeuMetTyrIleArgGlyAlaThrSerValGlnCysSerCysCysHisThrVal
 90 100

AsnLeuAlaLeuGluAlaAsnGlnValAlaHisValAsnCysGlyAsnCysMetMetLeu
 110 120

LeuMetTyrGlnTyrGlyAlaArgSerValLysCysAlaValCysAsnPheValThrSer
 130 140

ValGlyGlySerThrSerThrThrAspSerLysPheAsnAsn
 150

SEQ ID NO: 49

CysSerGlyCysArgAsnLeuLeuMetTyrProValGlyAlaThrSerValCysCysAlaValCys

SEQ ID NO: 50

CysGlyGlyCysHisThrLeuLeuMetTyrIleArgGlyAlaThrSerValGlnCysSerCysCys

SEQ ID NO: 51

CysGlyAsnCysMetMetLeuLeuMetTyrGlnTyrGlyAlaArgSerValLysCysAlaValCys

SEQ ID NO: 52

CysXxxXxxCysArgXxxXxxLeuMetTyrXxxXxxGlyAlaSerXxxValXxxCysXxxXxxCys

SEQ ID NO: 53

CysXxxXxxCysXxxXxxLeuLeuMetTyrXxxXxxGlyAlaXxxSerValXxxCysXxxXxxCys

SEQ ID NO: 54

10 30 50
 GAGGAGGAAGAGGAAGGTCCGCCCTCCGGGATGGGAATCTGCAGTTCTCCTCCTCCAATC
 70 90 110
 GTCACCATCACCGCCCGTAAACCCCAATCCCACCACCGTAGAAATTCCCGAAAAGGCC

130 150 170
 CAAATGGTATGTGGATCTTGCAGGCCTTGCTTCCTATCTAAGAGGATCCAAACATGTT
 190 210 230
 AAGTGCTCCTCTTGTCAAGACTGTTAACCTCGTTCTTGAAGCTAACCCAGGTTGGTCAGGTG
 250 270 290
 AATTGCAACAATTGCAAACGTCACTGATGTATCCTTATGGAGCTCCAGCTGTTAGATGT
 310 330 350
 TCCTCCTGCAATTCTGTCAACAGATATCAGTGAAAACAACAAACGACCTCCATGGTCTGAG
 370 390 410
 CAGCAAGGACCCTCAAAAGTTAACGCAGTCTCAGGAGAGCAGAGAATTAAACTTGAACC
 430 450 470
 GATTTTTGTCAATTGAAACCGTTTGACGACTAAAAACCTTGTAAATAATGTCGAAGGAT
 490
 AGATGAAATAAAATCACACC

SEQ ID NO:55

55

SEQ ID NO:56

CGSCRRLLSYLRGSKHVKCSSC

SEQ ID NO:57

CNNCKLLLMPYGAPAVRCSSC

SEQ ID NO: 58

10	30	50
GGAAGAGATACAACAAACAAACGCAGAAGGAAGAACAAAAGCACCGTGAAGAAGAAGAGGA		
70	90	110
GGAAGAGGAAGGTCCGCCCTCCGGGATGGGAATCTGCAGTTCTCCTCCTCCAATCGTCAC		
130	150	170
CATCACCGCCGCCGTAAACCCCAATCCCACCACCGTAGAAATTCCGAAAAGGCCAAAT		
190	210	230
GGTATGTGGATCTTGCAAGCGTTGCTTCTTATCTAAGAGGATCCAAACATGTTAAGTG		
250	270	290
CTCCTCTTGTCAAGACTGTTAATCTCGTTCTGAAGCTAACCAAGGTTGGTCAGGTGAATTG		
310	330	350
CAACAATTGCAAAGTGTACTGATGTATCCTTATGGAGCTCCAGCTGTTAGATGTTCC		
370	390	410
CTGCAATTCTGTCAACAGATATCAGTGTATGTATTCACAGATGGTTTGCTCCATGTCC		
430	450	470
AAATCATACTTGGAAAGAGTTGATACTTTGAGATCCGAGTAAGTAATCATCTGATGAAT		
490	510	530
CATTATAATAAAACTGTGTTATATTCAGGAAAACAACAAACGACCTCCATGGTCTGAGC		
550	570	590
AGCAAGGACCACTCAAAGTTAAGCAGTCTCAGGAGAGCAGAGAATTAAACCTGAACCG		
610	630	650
ATTTTTGTCAATTTGAACCGTTTGACGACTAAAAACCTTGTAATAATGTCGAAGGATA		
670	690	
GATGAAATAAAATCACCATTATAATCTAAAAAAAAAAAAAA		

SEQ ID NO:59

10	30	50
CTCTATCCTTACTCAACGGAGCTTACCAAGACCCAAACTCTCTAGGCCGACCGAGAG		

70 90 110
 TTGTTTGTACGTGTGCTTAACGCAGATTACATATGACGCTTCTAACCCACAATTAAATTG
 130 150 170
 GTTCACTCTTGCGCAAACCAAATAGCTAAAAAAGATTTAACCCAAATTCAATCC
 190 210 230
 TAAATCTGCATCATGGTCGGATAGTGTAGTGGCTGTGGCCTAATATCTACGCTATTGG
 250 270 290
 GGGATTCACTAATAAGAACCTTCGTCTAGCGTCATGGTCATGGATTGTCGTAC
 310 330 350
 TCACACATGGTGTAGGCCCTAGCATGCAGGTTCCCGTGTGTCATCTACTTGCCT
 370 390 410
 CCTTGATGGAAAATATATGTAACAGGAGGCCGCGGAACCTCGATTCAACGAAATGGAT
 430 450 470
 GGAGGTTTTGATACGAAAACCAAACCTGGAGTTTGCAATTCCCGAGTGAGGAGAA
 490 510 530
 GATATGCACAGGCTATAAGTGTGAGAGCATAGTGTATGAAGGAACGTCTATGTAAGGTC
 550 570 590
 GTATTTCTATAATGTGACTTACAAGCTGCATAAAGGTAGATGGATTCAAGCAGACTTT
 610 630 650
 AGGCGATGAATAATGGATGGCGTTGCTCATCTTTGTGTGATAAAGAACGTGTT
 670 690 710
 TACTTGTGCAATAGAAGTGGTAACGGTATGATCGATTGGTATGACTCGGAAAAGGATC
 730 750 770
 ATGGACAACATGAAGGGGTTGGAAAGATTGCCTAAAGTTATGGTAATGTTAAATTGGC
 790 810 830
 ATATTATGGTGGAAAAATGGTGGTGGTGTACGTGGAGTGCTAAGGAGTGGGTAAACGTGA
 850 870 890
 GAAAAAATTTGGTGTGCGGAAATTACGATTGAAAAACGCAAGGATGGAGAGATTGGGGA
 910 930 950
 TACTAGAAATGGTTGACGATGTATATAAAGCCAAGGATGAGCTAGAATATTTAGCTGTAG
 970 990 1010
 TGCATGCTGTTACTACCATCTGATTGATAAGAGAGTCATGTGAACATTGTCATTGA
 1030 1050 1070
 TTCACCGATGCAATAACGAATTATCTACTATCATTGTTGATTTCTTCTAAATCT
 1090 1110 1130
 TTTTGTTGTTCTGTATTGAATTACCTTACATTATTAAGAAAGTCAACTATTTGT

1150 1170 1190
 CAACGTTACTGGAAAGTTAAAAAGGTAAAAGTAATAATAATCTGAGAGTTAACCTTGGAC

 1210 1230 1250
 ATCTTCGCCGGAGCCGAGACGGAAGGCCTGATGGAAGAGATAACAACAAACGCGAGAAG

 1270 1290 1310
 GAAGAACAAAAGCACCGTGAAGAAGAAGAGGAGGAAGAGGAAGGTCCGCCTCCGGGATGG

 1330 1350 1370
 GAATCTGCAGTTCTCCTCCAATCGTCACCACCATCACCGCCCGTAAACCCCAATCCC

 1390 1410 1430
 ACCACCGTAGAAATTCCCGTATTCTGTAGTCTGTCTATTAGGGTTATCGATTTG

 1450 1470 1490
 CTTCCATTTCTTGCTACAGTCGATCAAATTAGAGATTTAGTGGAGTTGTAGACTTT

 1510 1530 1550
 TAGAGATAACCCATTTCGATTCCGAGAATTGATTAGTGTGTTTTCTGCAAATCTTCT

 1570 1590 1610
 TTGTTTTGGGTTGTTGCAGAAAAGGCCAAATGGTATGTGGATCTTGCAAGCGTTGC

 1630 1650 1670
 TTTCTTATCTAAGAGGATCCAAACATGTTAAGTGCCTCTGTCAAGACTGTTAATCTCG

 1690 1710 1730
 TTCTTGAAAGGTTCGTTCTCCATGGTTTTATCTCTTATTCAATTACTTGAAAAGCTTT

 1750 1770 1790
 TGTTGATAATCTCAGTCACTTGAAACTCTTAATGGAACAATCTTGGAAATGCTCTCAGT

 1810 1830 1850
 CTAGTTTACTTAGCATGTTGAATGATATATCTATGTTTTGAGAATCTCAAAATGT

 1870 1890 1910
 AAGCTTCCTGAGACCAATGAGTTAGTTCTTAACTGACACAAGAATGATCTGGTTAG

 1930 1950 1970
 GATTCTCTCTTAAGCTTTGTGAGCCTTTGGTCTCTACTCCATCATAATGTCTCCTT

 1990 2010 2030
 GTAGACCATTATGTTCTTACTCTTACTCTTACTCTTGGGGAAATTGTGTGAT

 2050 2070 2090
 CTTAAGACCAAGATTGTTCTTAGCTTGTGAATCACTTGGCCTCATTATTGATGAAAT

 2110 2130 2150
 AGCCTTCTCTCTTATCGGTTCTGGACTTGTCTTGTCTTGCAGCTAACCAAGGTTGG

 2170 2190 2210
 TCAGGTGAATTGCAACAATTGCAAACACTGCTACTGATGTATCCTTATGGAGCTCCAGCTGT

 2230 2250 2270

TAGATGTTCCCTCGCAATTCTGTACAGATATCAGTGTATGTATTACAGATGGTTTG
 2290 2310 2330
 TGCTCCATGTCCAATCATACTTGGAAAGAGTTGATACATTTGAGATCCGAGTAAGTAAT
 2350 2370 2390
 CATCTGATGAATCATTTATAATAAACTGTGTTATATTTCAGGAAAACAACAAACGACCTC
 2410 2430 2450
 CATGGTCTGAGCAGCAAGGACCCTCAAAAGTTAACGAGTCTCAGGAGAGCAGAGAATT
 2470 2490 2510
 AAACTTGAACCGATTTTGTCAATTTGAACCGGTTGACGACTAAAAACCTTGTAAATAA
 2530 2550 2570
 TGTCGAAGGATAGATGAAATAAAATCACCATTAATAATCTCATTGAATTCCCATTCTTC
 2590 2610 2630
 AGATATTACTTGTCAATCATCCTTACTGTTTAAGCTTAGTGGTTAAAAGAAATGTGT
 2650 2670 2690
 ATATATCCATACAAAAGTTGATATATGTACTGGACCAATATAACAAACACAGCTCACAG
 2710 2730 2750
 TCTCACACAATACATAAAACAAAATTCAATTTCACAGGTGAGAAAAACTAACTAGTAG
 2770 2790 2810
 TCTACTTGGCGAATTGTCAATGAATTCAATAATTAGTCGTATAATAGCAAACAAA
 2830 2850 2870
 ACATGGACTCTTACCCACCACAAATATGCATAATAATTACATTACAGTTCATATAAAA
 2890 2910 2930
 TACAAACTAATGGTGGGTCTCGAGAGAGCTAACAAAGAGCTGTGTGGGTGAAGAACCA
 2950 2970 2990
 ACTTGTCACGAAACCAATTAAATGAAATCAACCCCTAAATTAAATGAAACCTGGACGA
 3010 3030 3050
 AACTTACATTTGTTAACAGTTATCCTTTAAATCAAACCTGCATAGAATTGATTT

SEQ ID NO:60

MetGluGluIleGlnGlnGlnThrGlnLysGluGluGlnLysHisArgGluGluGluGlu
 10 20
 GluGluGluGluGlyProProProGlyTrpGluSerAlaValLeuProProProIleVal
 30 40
 ThrIleThrAlaAlaValAsnProAsnProThrThrValGluIleProGluLysAlaGln
 50 60
 MetValCysGlySerCysArgArgLeuLeuSerTyrLeuArgGlySerLysHisValLys
 70 80

SEQ ID NO:61

SEQ ID NO:62

CysXxxXxxCysXxxXxxLeuLeuXxxTyrXxxXxxGlyXxxXxxXxxValXxxCysSerSerCys

SEQ ID NO: 63

LeuValCysHisGlyCysArgAsnLeuLeuMetTyrProArgGlyAlaSerAsnValArgCysAlaLeuCysA
snThrIleAsnMetVal
IleIleCysGlyGlyCysArgThrMetLeuMetTyrThrArgGlyAlaSerSerValArgCysSerCysCysG
lnThrThrAsnLeuVal

IleAsnCysGlyHisCysArgThrThrLeuMetTyrProTyrGlyAlaSerSerValLysCysAlaValCysG
InPheValThrAsnVal

SEQ ID NO: 64

LeuValCysSerGlyCysArgAsnLeuLeuMetTyrProValGlyAlaThrSerValCysCysAlaValCysA
snAlaValThrAlaVal
LeuValCysGlyGlyCysHisThrLeuLeuMetTyrIleArgGlyAlaThrSerValGlnCysSerCysCysH
isThrValAsnLeuAla
ValAsnCysGlyAsnCysMetMetLeuLeuMetTyrGlnTyrGlyAlaArgSerValLysCysAlaValCysA
snPheValThrSerVal

SEQ ID NO: 65

MetValCysGlySerCysArgArgLeuLeuSerTyrLeuArgGlySerLysHisValLysCysSerSerCysG
InThrValAsnLeuVal
ValAsnCysAsnAsnCysLysLeuLeuLeuMetTyrProTyrGlyAlaProAlaValArgCysSerSerCysA
snSerValThrAspIle

SEQ ID NO: 66

Nucleic acid sequence of C

10	30	50
AGCAACAACAACAACCAGCAACCACCAACCTCCGTCTATCCACCTGGCTCCGCC		
70	90	110
GTCACAACCGTAATCCCTCCTCCACCATCTGGATCTGCATCAATAGTCACCGGAGGAGGA		
130	150	170
GCGACATACCACCATCCTCCAGCAACACAGCAACAGCTCAAATGTTCTGGACATAAC		
190	210	230
CAGAGACAAGAGATCGAACAGGTAAACGATTTCAAAAACCATCAGCTCCCTCTAGCTCGT		
250	270	290
ATCAAAAAATCATGAAAGCTGATGAAGATGTGCGTATGATCTCCGCCGAAGCACCGATT		
310	330	350
CTCTTCGCGAAAGCTTGTGAGCTTTCAATTCTCGAACTTACGATTAGATCTGGCTTCAC		
370	390	410
GCTGAAGAGAACAAACGTCGTACGCTTCAGAAAAACGATATCGCTGCTGCGATTACTAGA		
430	450	470
ACCGATATCTCGATTTCTTGTGATATTGTTCTAGGGAAGAGATCAAGGAAGAGGAA		
490	510	530
GATGCAGCATCGGCTCTGGTGGAGGAGGTATGGTTGCTCCGCCGAGCGGGTGTTCCT		

550	570	590
TATTATTATCCACCGATGGGACAACCGGGCGGTTCCCTGGAGGGATGATGATTGAAAGACCG		
610	630	650
GCGATGGATCCTAGCGGTGTTATGCTCAGCCTCCTCTCAGGCATGGCAAAGCGTTGG		
670	690	710
CAGAATTCACTGGTGGTGATGATGTGTCCTATGGAAGTGGAGGAAGTAGCGGCCAT		
730	750	770
GGTAATCTCGATAGCCAAGGTTGAGCTATGGAACCAGAACGCTTAGAGATTAAATCATCAT		
790	810	830
TTCGACCCCTGCAAGTGTCTGATTCTTATGCTATGATTCAATGACTTA		

SEQ ID NO: 67

Amino acid sequence of C

SerAsnAsnAsnAsnAsnGlnGlnProProProThrSerValTyrProProGlySerAla	
10	20
ValThrThrValIleProProProSerGlySerAlaSerIleValThrGlyGlyGly	
30	40
AlaThrTyrHisHisLeuLeuGlnGlnGlnGlnGlnLeuGlnMetPheTrpThrTyr	
50	60
GlnArgGlnGluIleGluGlnValAsnAspPheLysAsnHisGlnLeuProLeuAlaArg	
70	80
IleLysLysIleMetLysAlaAspGluAspValArgMetIleSerAlaGluAlaProIle	
90	100
LeuPheAlaLysAlaCysGluLeuPheIleLeuGluLeuThrIleArgSerTrpLeuHis	
110	120
AlaGluGluAsnLysArgArgThrLeuGlnLysAsnAspIleAlaAlaAlaIleThrArg	
130	140
ThrAspIlePheAspPheLeuValAspIleValProArgGluGluIleLysGluGluGlu	
150	160
AspAlaAlaSerAlaLeuGlyGlyGlyMetValAlaProAlaAlaSerGlyValPro	
170	180
TyrTyrTyrProProMetGlyGlnProAlaValProGlyGlyMetMetIleGlyArgPro	
190	200
AlaMetAspProSerGlyValTyrAlaGlnProProSerGlnAlaTrpGlnSerValTrp	
210	220

GlnAsnSerAlaGlyGlyAspAspValSerTyrGlySerGlyGlySerSerGlyHis
 230 240
 GlyAsnLeuAspSerGlnGly

SEQ ID NO: 68**Nucleic acid sequence of CC**

10 30 50
 AGTATGGATGAGCTTCAGAACGCTCTCAGATACTCACATGTTGCTTGACATGGTGTAC
 70 90 110
 TGCACGGTTTGCATGTATGCAGACACAACACAAGATGGAAATGGACAAGAGGGACGGT
 130 150 170
 AAGTTGGGCCACAGCCAATGGCAGTGCCTCCGGCTCAGCAAATGTCACGGTTGATCAA
 190 210 230
 GCCACCCCACCCGCAGTCGGTTATCCTCCACAACAAGGTTATCCACCTCTGGTTATCCT
 250 270 290
 CAACACCCCTCCACAAGGTTATCCACCTCTGGCTATCCTCAAAACCCCTCCCTCAGCT
 310 330 350
 TATTCTCAATACCCTCCTGGGGCTTATCCTCCTCCCGCTTACCCAAAGTGATCACTC
 370 390 410
 TTTGCCTGTTTCTCTCCGATTGGAAAATTTTATTCATCTTTTTAATGCTGTCTTG
 430 450 470
 TTACGGGTCAAGAATTGAACGTTCGCTGATTGTTTGAGGTCGTTGTTGATGAGATT
 490 510 530
 TGACCTCGCATGTTGTTGTTCTGAAACGTCCCTTGGACTAAGAGATTTCATGA
 550
 CTTAAAAAAAAAAAAAA

SEQ ID NO: 69**Amino acid sequence of CC**

SerMetAspGluLeuSerGluAlaSerGlnIleLeuThrCysCysSerAspMetValTyr
 10 20
 CysThrValCysAlaCysMetGlnThrGlnHisLysMetGluMetAspLysArgAspGly

30

40

LysPheGlyProGlnProMetAlaValProProAlaGlnGlnMetSerArgPheAspGln
 50 60

AlaThrProProAlaValGlyTyrProProGlnGlnGlyTyrProProSerGlyTyrPro
 70 80

GlnHisProProGlnGlyTyrProProSerGlyTyrProGlnAsnProProProSerAla
 90 100

TyrSerGlnTyrProProGlyAlaTyrProProProProAlaTyrProLys
 110

SEQ ID NO: 70**Nucleic acid sequence of FF**

10 30 50
 AGGTTTCCGACGTTGATGACCCAATTCCGTCGTCGACGAAGACGATTCCGGCATCGTAT

70 90 110
 TTGCTTCCGTTACAATGGCCTCAGCCGCAGAACGAGGAGATTCTCTGCCATGGAAGAA

130 150 170
 GCTGAGTTCGAAGAAAAGTGCACACGAGATCAGAAAGATGAGTCCTGCTTACCGTAATT

190 210 230
 GGAAAACCAGTCGTCAACAACGAACAAGAACGAGGATGATAATGAATCAGAGGATGATGAT

250 270 290
 GCAGATAATGCAGAGGAATCAGATGGTGAAGAGAGTTGAGCAAGAAACCGGATAAATAATC

310 330 350
 TTGAGGCCGAAAATACACAAGGGTTATTGATGGCATTGGCTTGAAACTTGAGGACCCCTA

370 390 410
 TCTAAATCTTCTTGTGATAAAACGACTGTGATTCTGACTTGTAAACCANGTTTTTCT

430 450 470
 TTTCTTAGGAACGACTGAAATGTTCACTTTGGCCCTAAGGTTAGTCAGTGGATTATCG

490 510 530
 TAGTTAATTGTCTCAATCTCATGGTGTAAATTGTGTTAGTGTATTGACATTGAATTTAT

550 570
 GGTTTATAGATTGTAGTGATTGATGAAAAA

SEQ ID NO: 71

Amino acid sequence of FF

SEQ ID NO: 72

Nucleic acid sequence of GG

10 30 50
 AGGGAAACAATGAGCCAGTACAATCAACCTCCC GTTGGT GTCCTCCTCAAGGTTAT
 70 90 110
 CCACCGGAGGGATATCCAAAAGATGCTTATCCACCACAAGGATATCCTCCTCAGGGATAT
 130 150 170
 CCTCAGCAAGGCTATCCACCTCAGGGATATCCTCAACAAGGTTATCCTCAGCAAGGATAT
 190 210 230
 CCTCCACCGTAGCGCCTCAATATCCTCCACCACCGCAGCATCAGCAACAACAGAGCAGT
 250 270 290
 CCTGGCTTCTAGAAGGATGTCTTGCTGCTCTGTGTTGCTGTCCTGGATGCTTGC
 310 330 350
 TTCTGATTGGAGTCTCTCTCTGCATAAAGCTTCGGGATTTATTTGTAAGAGGGTTT
 370 390 410
 TGGTTAACAAAAACCTTAATTGATTTGTGGGCATTAAAAATGAATCTCTCGATGATT
 430 450 470
 TCTTTCGTTTATGTGTAATGTTCTCGGGTCATAACATTAACTATTGTCTATCGACG
 490 510 530
 TTCTGCCTTAGTTGTATTTGATTATGGGAATGTAAATTGGTTGGGAGACACTATTCTAT

550 570
GCCATAGTTATTGCTTGGATCTTCAAAAAAAAAAAAAA

SEQ ID NO: 73

Amino acid sequence of GG

Arg	Glu	Thr	Met	Ser	Gln	Tyr	Asn	Gln	Pro	Pro	Val	Gly	Val	Pro	Pro	Pro	Gln	Gly	Tyr
10																		20	
Pro	Pro	Glu	Gly	Tyr	Pro	Lys	Asp	Ala	Tyr	Pro	Pro	Gln	Gly	Tyr	Pro	Pro	Gln	Gly	Tyr
30																		40	
Pro	Gln	Gln	Gly	Tyr	Pro	Pro	Gln	Gly	Tyr	Pro	Gln	Gln	Gly	Tyr	Pro	Gln	Gln	Gly	Tyr
50																		60	
Pro	Pro	Pro	Tyr	Ala	Pro	Gln	Tyr	Pro	Pro	Pro	Gln	His	Gln	Gln	Gln	Gln	Ser	Ser	
70																		80	
Pro	Gly	Phe	Leu	Glu	Gly	Cys	Leu	Ala	Ala	Leu	Cys	Cys	Cys	Cys	Leu	Leu	Asp	Ala	Cys
90																		100	

Phe

SEQ ID NO: 74

Nucleic acid sequence of HH

10	30	50
AGTGATGTTCTCCTAAGTCGGTTGACTGGAGAAACGAAGGCGCAGTGACTGAAGTCAAA		
70	90	110
GATCAAGGCCCTTGAGGAGTTGGGGCTTCTCCACTGTGGGAGCAGTGGAAAGGCTTA		
130	150	170
AACAAGATTGTGACTGGAGAGCTAGTAACCTTGTCTGAGCAAGATTGATCAATTGTAAC		
190	210	230
AAAGAAAACAATGGTTGCGGAGGAGGCAAAGTCGAGACAGCCTATGAGTCATCATGAAC		
250	270	290
AATGGTGGCTTGGTACCGACAACGATTATCCTTACAAAGCTCTCAATGGAGTCTGCGAA		
310	330	350
GGCCGCCTCAAGGAAGACAACAAGAATGTTATGATTGATGGGTATGAGAATTGCCTGCA		
370	390	410
AACGATGAAGCCGCTCTCATGAAAGCGGTTGCTCACCGCCTGTGACTGCCGTTGCGAT		

430 450 470
 TCCAGCAGCCGAGAGTTTCAGCTTATGAATCGGGAGTGTGACGAACTTGCGAACAA

 490 510 530
 AACCTAACCATGGTGTGTTGGTCGGGTATGGAACCGAGAATGGTCGTACTACTGG

 550 570 590
 ATTGTGAAAAACTCGAGGGGCACACATGGGGGGAGGCTGGCTACATGAAGATGGCTCGC

 610 630 650
 AACATTGCCAATCCAAGAGGCATATGTGGCATCGCAATGCGAGCTCATACCCCTCTCAAG

 670 690 710
 AACTCGTTTCTACGGATAAAGTTCGGTTGCCTAATAATATGAACAAATGTATGCCAT

 730 750 770
 GGAACGGATCGGTTAACCCATTATCGTTATTCGACTTGAAGGAAACTAAAAAATAATGT

 790 810 830
 GGTCGATTGGTTGGTTTGTATATATTATGCATTTGTATGGGGTCAGTCATGTTG

 850 870 890
 AACTTTGTATAATATTTCTTGGGTCTAGTGATAAATAATTTCCCTTGCAGAAAAAAA

 910
 AAAAAAAAAAA

SEQ ID NO: 75

Amino acid sequence of HH

SerAspValLeuProLysSerValAspTrpArgAsnGluGlyAlaValThrGluValLys
 10 20
 AspGlnGlyLeuCysArgSerCysTrpAlaPheSerThrValGlyAlaValGluGlyLeu
 30 40
 AsnLysIleValThrGlyGluLeuValThrLeuSerGluGlnAspLeuIleAsnCysAsn
 50 60
 LysGluAsnAsnGlyCysGlyGlyLysValGluThrAlaTyrGluPheIleMetAsn
 70 80
 AsnGlyGlyLeuGlyThrAspAsnAspTyrProTyrLysAlaLeuAsnGlyValCysGlu
 90 100
 GlyArgLeuLysGluAspAsnLysAsnValMetIleAspGlyTyrGluAsnLeuProAla
 110 120
 AsnAspGluAlaAlaLeuMetLysAlaValAlaHisGlnProValThrAlaValValAsp
 130 140

SerSerSerArgGluPheGlnLeuTyrGluSerGlyValPheAspGlyThrCysGlyThr
 150 160
 AsnLeuAsnHisGlyValValValValGlyTyrGlyThrGluAsnGlyArgAspTyrTrp
 170 180
 IleValLysAsnSerArgGlyAspThrTrpGlyGluAlaGlyTyrMetLysMetAlaArg
 190 200
 AsnIleAlaAsnProArgGlyIleCysGlyIleAlaMetArgAlaSerTyrProLeuLys
 210 220
 AsnSerPheSerThrAspLysValSerValAla
 230

SEQ ID NO: 76**Nucleic acid sequence of I**

10 30 50
 AGCGAAATGCCAGTTCAAGCTCCATCTCCGCCTCGTCTTCATTCTCCGTTCAATTCACTGT
 70 90 110
 CCCATCAATTCACTCCTTCTTCTCGGCGAGGAATCTCCGGTCGCCGTCAACATCT
 130 150 170
 TATCCCCGAATCAAAGCTGAACCTCGATCCAACACCGTAGTCGCGATATCTGTAGGCGTA
 190 210 230
 GCAAGCGTCGCATTAGGAATCGGAATCCCTGTGTTCTACGAGACTCAAATCGACAATGCG
 250 270 290
 GCTAAGCGAGAGAAATACTCAACCTTGT'TTCCCTGTAATGGCACCGGAGCTCAGAAATGC
 310 330 350
 AGATTGTGTGGAAAGTGGTAATGTGACCGTAGAGCTTGGTGGAGGAGAGAAAGAAGTC
 370 390 410
 TCAAAACTGTATCAACTGTGATGGTGTGGTCTTAACTTGCACTACTTGTCAAGGCTCT
 430 450 470
 GGTGTTCAACCTCGATACTTGATCGAAGGGAGTTCAAGGACGATGACTAAATACCTTGC
 490 510 530
 TCTAAGGAACATTTCTTTCTTCTCCCTTCTCACATTCTCATTGTACAATGCTGTTT
 550 570 590
 GTTCACCAAAACATGTTGAGAGAACATCATGACATGGATATTGTAATTGTGAAAGAAAACC
 610 630 650

ACCAGAGTTCAATCAAATGTTCTTCTTGACTTAAAAAAAAAAAAAA

SEQ ID NO: 77

Amino acid sequence of I

Ser	Glu	Met	Pro	Val	Ser	Ala	Pro	Ser	Pro	Pro	Arg	Leu	His	Ser	Pro	Phe	Ile	His	Cys
10																			20
Pro	Ile	Asn	Phe	Thr	Pro	Ser	Ser	Phe	Ser	Ala	Arg	Asn	Leu	Arg	Ser	Pro	Ser	Thr	Ser
	30																	40	
Tyr	Pro	Arg	Ile	Lys	Ala	Glu	Leu	Asp	Pro	Asn	Thr	Val	Val	Ala	Ile	Ser	Val	Gly	Val
																		50	
Ala	Ser	Val	Ala	Leu	Gly	Ile	Gly	Ile	Pro	Val	Phe	Tyr	Glu	Thr	Gln	Ile	Asp	Asn	Ala
																		70	
Ala	Lys	Arg	Glu	Asn	Thr	Gln	Pro	Cys	Phe	Pro	Cys	Asn	Gly	Thr	Gly	Ala	Gln	Lys	Cys
																		90	
Arg	Leu	Cys	Val	Gly	Ser	Gly	Asn	Val	Thr	Val	Glu	Leu	Gly	Gly	Glu	Lys	Glu	Val	
																		110	
Ser	Asn	Cys	Ile	Asn	Cys	Asp	Gly	Ala	Gly	Ser	Leu	Thr	Cys	Thr	Thr	Cys	Gln	Gly	Ser
																		130	
Gly	Val	Gln	Pro	Arg	Tyr	Leu	Asp	Arg	Arg	Glu	Phe	Lys	Asp	Asp	Asp				
																		150	

SEQ ID NO: 78

Nucleic acid sequence of II

10	30	50
AGAGAAAACATGGGAGGTGACAATGATAATGACAAAGACAAAGGGTTCATGGTATCCT		
70	90	110
CCCGCTGGATACCCACCCCCTGGGGCTTATCCACCCGCTGGATACCCACAACAAGGTTAC		
130	150	170
CCTCCACCACCCGGTGCTTACCCGCCTGCAGGTTATCCTCCGGGTGCCTACCCACCTGCT		
190	210	230
CCTGGTGGTTATCCTCCGCCCTGGTTATGGTGGTTATCCTCCAGCTCCTGGTTATGGA		
250	270	290

GGTTATCCTCCTGCACCTGGTCATGGTGGTACCCCTCCTGCTGGCTATCCTGCTCATCAC
 310 330 350
 TCAGGACACGCAGGAGGAATTGGGGGTATGATTGCAGGTGCTGCAGCTGCCTATGGAGCT
 370 390 410
 CACCACGTATCTCATAGCTCTCACTGTCCTAACGGACATGCTGCATATGGTCACGGTTTT
 430 450 470
 GGCCATGGTCATGGCTATGGCTATGGTCATGGTAAGTTCAAGCATGGAAAGCAC
 490 510 530
 GGGAAAGTTCAAGCATGGAAAGCATGGAATGTTGGAGGAGGCAAGTTCAAGAAGTGGAAAG
 550 570 590
 TGATCTAGCTATTACCTTGTGAATTGTCTGGACTGACCAATGTTCAAATAAGCCCT
 610 630 650
 AAACATTATATAAGTTGACTTCGTCGGTAGATTGCTGGTCGAGTTGGAATAATTGAA
 670 690 710
 ACTTAATTAGTATCAAATCTTATTGTGTACTTTAAAGCTATCGTGGCTTATAATGACA
 730 750 770
 GATTCTGGTTTCGGTGTGTTAGATTTGTATATACTGTTTTACATTGCTTA
 790 810
 AGCTTATAGAAGTCATGATTATGATTAAAAAAAAAAAAAAA

SEQ ID NO: 79

Amino acid sequence of II

ArgGluAsnMetGlyGlyAspAsnAspAsnAspLysAspLysGlyPheHisGlyTyrPro
 10 20
 ProAlaGlyTyrProProProGlyAlaTyrProProAlaGlyTyrProGlnGlnGlyTyr
 30 40
 ProProProProGlyAlaTyrProProAlaGlyTyrProProGlyAlaTyrProProAla
 50 60
 ProGlyGlyTyrProProAlaProGlyTyrGlyGlyTyrProProAlaProGlyTyrGly
 70 80
 GlyTyrProProAlaProGlyHisGlyGlyTyrProProAlaGlyTyrProAlaHisHis
 90 100
 SerGlyHisAlaGlyGlyIleGlyGlyMetIleAlaGlyAlaAlaAlaAlaTyrGlyAla
 110 120
 HisHisValSerHisSerSerHisCysProTyrGlyHisAlaAlaTyrGlyHisGlyPhe

130

140

GlyHisGlyHisGlyTyrGlyTyrGlyHisGlyHisGlyLysPheLysHisGlyLysHis
 150 160

GlyLysPheLysHisGlyLysHisGlyMetPheGlyGlyGlyLysPheLysLysTrpLys
 170 180

SEQ ID NO: 80**Nucleic acid sequence of K**

10	30	50
AGTGTCACTACTCCATCCGAGGGAGGATTCAAACAAACGGTTTACCGGTTTCAGCAACCCGGT		
70	90	110
ACACCGAACCAACCAGCGAACCAAGAGTTCCCGTAGTCATTGCGCCGCCGAATTATCAGCAA		
130	150	170
GCTAATGTTAACCTATCTGTTGGGAGGCCATGGAGCAGTGGTTGTTGATTGTCAAGCA		
190	210	230
GACCAAGCCAATGCCGTTTGACCACAATTGTACCTTGTAAACATTGGACAAATAGCA		
250	270	290
GAAGTGATGGATGAAGGAGAGATGACTTGTCCCTTTGGAACTTTCATGTACTTATTGATG		
310	330	350
ATGCCGGCTTTATGCTCTCACTGGGTGATGGGATCAAAGTATAGAGAAAAAAATGAGGAGA		
370	390	410
AAATTAAATCTTGTGGAAGCTCCATATTCAAGATTGTGCCAGTCATGTCCATTGCCCTTGT		
430	450	470
TGCTCTTTGTCAAGAACATACAGAGAGCTCAAGATTAGGAATCTTGATCCTCTCTAGGT		
490	510	530
TGGAATGGGATACTTGCTCAAGGACAAGGACAATATGAGAGAGAGAACACCAAGTTTGCT		
550	570	590
CCTACAAATCAATATATGTCTAACGAAACATTGATTAGTTGACTTCCATATTATTA		
610	630	650
AAACATTATTGTGGACCATTGTACAATGAAAGTGTGCTATATTAAAATTGCAATGCAA		
670	690	
GTGTGAGATTGATAAAAAAAAAAAAAAA		

SEQ ID NO: 81

Amino acid sequence of K

SerValThrThrProSerGluGluAspSerAsnAsnGlyLeuProValGlnGlnProGly
10 20

ThrProAsnGlnArgThrArgValProValSerGlnPheAlaProProAsnTyrGlnGln
30 40

AlaAsnValAsnLeuSerValGlyArgProTrpSerThrGlyLeuPheAspCysGlnAla
50 60

AspGlnAlaAsnAlaValLeuThrThrIleValProCysValThrPheGlyGlnIleAla
70 80

GluValMetAspGluGlyGluMetThrCysProLeuGlyThrPheMetTyrLeuLeuMet
90 100

MetProAlaLeuCysSerHisTrpValMetGlySerLysTyrArgGluLysMetArgArg
110 120

LysPheAsnLeuValGluAlaProTyrSerAspCysAlaSerHisValLeuCysProCys
130 140

CysSerLeuCysGlnGluTyrArgGluLeuLysIleArgAsnLeuAspProSerLeuGly
150 160

TrpAsnGlyIleLeuAlaGlnGlyGlnGlyGlnTyrGluArgGluAlaProSerPheAla
170 180

ProThrAsnGlnTyrMetSerLys

SEQ ID NO: 82

Nucleic acid sequence of M

10 30 50
AGAAAATACGAAAAGGTCTCCCTCCCAGCACCTAACGTGGCTGGACACTCGAGCCATCAC

70 90 110
GAAGACGACGGTCAATACTATCCCGGCAAATACGAAAAAGCCTCCCTCCCAGCACCTAC

130 150 170
GTGGCCGGATATCCGAGCCATCATGAAGACGATGGTCAATACTATCCTGGCAAATACGAA

190 210 230
AAGGTCTCCCTCCCAGCACCTAACGTGGCTGGACACCCGAGCCACTCCGAAGATGATGGC

250 270 290
CAATACTATCCCGGCAAATACGAAAAGGCCTCCGTCCCATCAGCTTACGTGGCCGAACAC

310 330 350
TCGAGCCACTCCGAAGATGATGGCCAATACTATCCTGGCAAATACGAAAAGCCCGAACAC

370 390 410
 CATTACTGAAAACCTCTCACACACAATGATTCTCATCCTTCCGTAGCTTTAATTGAC
 430 450 470
 TTTTAACAATAAAAACGTATCTTAATTTTCATCAAAAAAAAAAAAAAA

SEQ ID NO: 83

Amino acid sequence of M

SEQ ID NO: 84

Nucleic acid sequence of Oo

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          10          30          50
AGCCGATCTCAGATTCTTCCATCTTCCAGGAGGAATTCAGTGTGGCGACCACACAGCTT

          70          90          110
GGCATTCCAACAGACGATCTAGTCGGCAATCACACCGCCAAATGGATGCGAGGATAGAAGC

          130         150         170
AAGAAATCACCTATGGAACTGATTAGTAGTGAGGTTCCACCTATCAAAGTTGATGGAAGGATT

          190         210         230
GTTGCTTGTGAAGGAGACACCAATCCGGCCCTAGGTCATCCAATCGAGTTCATATGCCTC

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250	270	290
GACCTAAATGAGCCTGCGATCTGCAAGTACTGCGGCCCTCGTTATGTTCAAGATCATCAC		
310	330	350
CATTGAGGCAAATTCTGAAAGTGAAC TGCTGGTCTCTCTCCCTTTATTGCATTTTA		
370	390	410
AGTTTGTTGTATTGTTTTCTGGTGTGCCTACTACATCTTCAGCTATATTATCTAATAA		
430	450	470
AGGATT CGATCAAAGTCGGGTAAGTTGATTTGATCTCAC TT CAGCACTTGTC		
490	510	530
TGTTGTAACATTCAATCTCTGATATCACTGTC TTTACATGCCAAAAAAAAAAAAAA		
550 AAAAAAAAAAAAAAA		

SEQ ID NO: 85**Amino acid sequence of O0**

SerArgSerGlnIleLeuProSerSerArgArgAsnPheSerValAlaThrThrGlnLeu	20
10	
GlyIleProThrAspAspLeuValGlyAsnHisThrAlaLysTrpMetGlnAspArgSer	40
30	
LysLysSerProMetGluLeuIleSerGluValProProIleLysValAspGlyArgIle	60
50	
ValAlaCysGluGlyAspThrAsnProAlaLeuGlyHisProIleGluPheIleCysLeu	80
70	
AspLeuAsnGluProAlaIleCysLysTyrCysGlyLeuArgTyrValGlnAspHisHis	100
90	
HisEndGlyLysPhe	

SEQ ID NO: 86**Nucleic acid sequence of P**

10	30	50
AGAACAGCTCGAGTTCTTATGGGCCTAGACTCTCTGGTGGTACAACCGATCTGGA		
70	90	110
AACAGGGTTCCCGCGTAACAAACCAAGCTTCCCAATAGCACCAGTCCAATGGTGAGGCT		

130 150 170
 AATCAATTCAATGGCCCAAGAATAATGAACCCCCATGCTGCTGAGTTCATACCGAGTCAA
 190 210 230
 CCTTGGGTTCTAATGGGTATCCAGTGTCAACCAATGGCTATTTAGCATCCCCAAATGGT
 250 270 290
 GCAGAAAATAACACAGAATGGGTACCCCTTGTCAACCAGTAGCAGGTGGATATCCGTGTAAC
 310 330 350
 ATGTCCGTTACACAGCCTCAGGATGGACTTGTTCAGAGGAATTACCTGGTGCTGGAAC
 370 390 410
 TCTGAGGAGAAGAGCGGAAGCGAAGAAGAAAGCAACAAACGACAAAAATGCTGGAGAGGAT
 430 450 470
 GACGAAGCCGTTGGACAAGAAACTACAGATACACCTGAAAATGGACATTGACAGTAGGT
 490 510 530
 GAAGTGGAAACCCACATCACATGAGACTTGTGATGAGAAAAATGGAGAACGACAAGGAGC
 550 570 590
 AAGTGCTGGGAGATTACAGCGATAATGAAATCGAGCAAATTGAAGTTACAAGTTGAAGA
 610 630 650
 CGCAACTGTCTGTTACTGAAGTATTAACATTGAGGCTAAAGGAATGCGGAGACATTTGG
 670 690 710
 CTCCATTGATGAGGTTAAAGGTAAACAATCATCATAGTCGAGAAAAGCATTTCATGTT
 730 750 770
 GAATGTTTGTGTTAGCGCAGGACCAAGGCTGTCACTCCTGTTAACAACTTTCT
 790 810 830
 CCTGCTTTCAGTTTGGTTCATAGCTGAAACTAGATATTCACACTCCTTAACAAAA
 850 870
 GATTTGTCCCTTGTGTTAAAAAAAAAAAAAAAAAAAAAA

SEQ ID NO: 87

Amino acid sequence of P

35

67

70	80
AlaGluIleThrGlnAsnGlyTyrProLeuSerProValAlaGlyGlyTyrProCysAsn	
90	
MetSerValThrGlnProGlnAspGlyLeuValSerGluGluLeuProGlyAlaGlySer	
110	
SerGluGluLysSerGlySerGluGluGluSerAsnAsnAspLysAsnAlaGlyGluAsp	
130	
AspGluAlaValGlyGlnGluThrThrAspThrProGluAsnGlyHisSerThrValGly	
150	
GluValGluThrThrSerHisGluThrCysAspGluLysAsnGlyGluArgGlnGlyGly	
170	
LysCysTrpGlyAspTyrSerAspAsnGluIleGluGlnIleGluValThrSer	
190	

SEQ ID NO: 88

Nucleic acid sequence of T

10	30	50
AGAGACCATCCAGCTTACCATCAGATCCACCAGCAACAACAACAGCTCACTAACAG		
70	90	110
CTTCAATCTTTCTGGGAGACTCAATTCAAAGAGATTGAGAAAACCACTGATTCAAGAAC		
130	150	170
CATAGCCTTCATTGGCAAGAATCAAGAAAATCATGAAAGCTGATGAAGATGTGGTATG		
190	210	230
ATCTCGGCCGAGGCCTGTTGTTGCCAGGGCTGCGAGATGTTATTCTGGAGCTT		
250	270	290
ACGTTAACGGCTTGAAACCATACTGAGGAGAACAGAGAAGGACGTTGCAGAACATGAT		
310	330	350
ATCGCGGCTGCGGTGACTAGAACTGATATTTTGATTTCTGTGGATATTGTTCTCGG		
370	390	410
GAGGATCTCGTATGAAAGTCTGGGTGGTGGTGTGAAGCTGCTACAGCTGCGGGT		
430	450	470
TATCCGTATGGATACTTGCCTCTGGAACAGCTCCAATTGGAACCCGGGAATGGTTATG		
490	510	530
GGTAACCCGGCGCGTATCCGCCGAACCCGTATATGGTCAGCCAATGTGGCAACAAACCA		
550	570	590
GGACCTGAGCAGCAGGATCCTGACAATTAGCTTGGCTAATAACTAGCCGTCTAATTG		

610 630 650
AAGCTCTCCCCGGTGGATCTACTCAAGAAGAAGAATGTTAATAGAAAACATTGCGACAT

670 690 710
AAAAAGTTGGTAGTAGAATAATTCTGTTTATGATCCATGGATTATCTATTGTTA

730 750 770
TTCAGTTGGTTATCTGTCATCAAACGTGTTTGGTCAATGTAACAAATTCAAAACT

790 810 830
GAGAATTGAACCTACAAAGGCTAGATTACTACTTATAAAGTTCAAAGCTAAAAAAAAAA

AAAAAAA

SEQ ID NO: 89

Amino acid sequence of T

ArgAspHisProAlaTyrHisGlnIleHisGlnGlnGlnGlnGlnGlnLeuThrGlnGln	20
10	
LeuGlnSerPheTrpGluThrGlnPheLysGluIleGluLysThrThrAspPheLysAsn	40
30	
HisSerLeuProLeuAlaArgIleLysLysIleMetLysAlaAspGluAspValArgMet	60
50	
IleSerAlaGluAlaProValValPheAlaArgAlaCysGluMetPheIleLeuGluLeu	80
70	
ThrLeuArgSerTrpAsnHisThrGluGluAsnLysArgArgThrLeuGlnLysAsnAsp	100
90	
IleAlaAlaAlaValThrArgThrAspIlePheAspPheLeuValAspIleValProArg	120
110	
GluAspLeuArgAspGluValLeuGlyGlyValGlyAlaGluAlaAlaThrAlaAlaGly	140
130	
TyrProTyrGlyTyrLeuProProGlyThrAlaProIleGlyAsnProGlyMetValMet	160
150	
GlyAsnProGlyAlaTyrProProAsnProTyrMetGlyGlnProMetTrpGlnGlnPro	180
170	
GlyProGluGlnGlnAspProAspAsn	

SEQ ID NO: 90**Nucleic acid sequence of X**

10	30	50
AGATT CGT ATT CCT GG CAA AGA AAG ACA AGA ATT CTG TT ACAG TGG ACT TCAG GAA ATC		
70	90	110
GAT GTG AACT CTG AGC TT GTT GT ATCC ACC GACT CTG CCC GACC ATT GGT GAAT ACT GAA		
130	150	170
GAT GTG AGA AAG GT CCT AA AGA TGG TCC CG CG TT GG AGC AGC TG ACT TT GGT GTT CCT		
190	210	230
GCT AA AGC TAC AA TCA AAG AGG TCA ATT CTG ATT CG CCT GT GGT GAA AAC TCT GAC AGA		
250	270	290
AAA ACC CT AT GGG AA AT GC AG AC ACC AC AG GT GAT CA AAC CC AG AG CT ATT GAA AAA AGG GT		
310	330	350
TT CG AGC TT GT AAA AGT GA AGG TCT AG AGG TA AC AG AT GA CG TT CG ATT GT TGA AT AC		
370	390	410
CT CA AGC AT CC AG TT AT GT CT CT CA AGG AT CT TAC AA AC AT CA AGG TT AC AA AC CCT		
430	450	470
GAT GAT T TACT GCT TGT GAG AGA AT CTT GAG CG AGG ACT CAT GAG AT ATT AT AT CATT		
490	510	530
ACT TAG T AAG AAG AC GT GT CA AGG GT AT GC AT GAA AA AT GT TT ATT GAA AT CTT GC AT		
550	570	590
CCT AG TT GGT GG TT AT AAA AT GT GCA AG AT AAT GT TT CACT GAA AAC TACT T GCT GT		
610	630	650
GA AT AT GG AT TC GA AC AG AG CC AAT CG AAG TGA AT TT GC AT ATT GT AAA A A A A A A A A		
670		
A A A A A A A A		

SEQ ID NO: 91**Amino acid sequence of X**

Arg Phe Ala Ile Pro Gly Lys Glu Arg Gln Asp Ser Val Tyr Ser Gly Leu Gln Glu Ile		
10		20
Asp Val Asn Ser Glu Leu Val Cys Ile His Asp Ser Ala Arg Pro Leu Val Asn Thr Glu		
30		40
Asp Val Glu Lys Val Leu Lys Asp Gly Ser Ala Val Gly Ala Ala Val Leu Gly Val Pro		
50		60

AlaLysAlaThrIleLysGluValAsnSerAspSerLeuValValLysThrLeuAspArg
70 80

LysThrLeuTrpGluMetGlnThrProGlnValIleLysProGluLeuLeuLysLysGly
90 100

PheGluLeuValLysSerGluGlyLeuGluValThrAspAspValSerIleValGluTyr
110 120

LeuLysHisProValTyrValSerGlnGlySerTyrThrAsnIleLysValThrThrPro
130 140

AspAspLeuLeuLeuAlaGluArgIleLeuSerGluAspSer
150

THE CLAIMSWhat is Claimed Is:

5

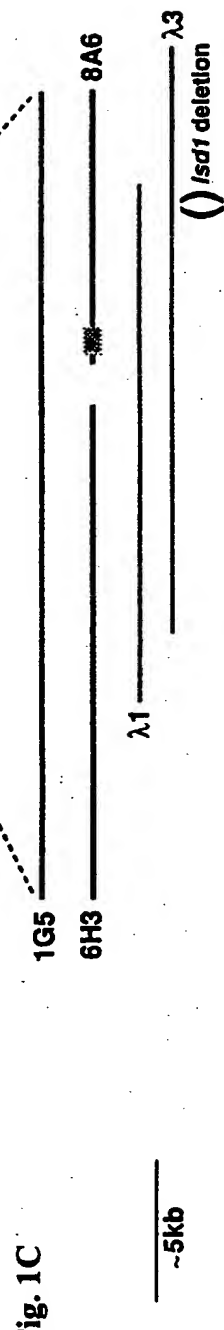
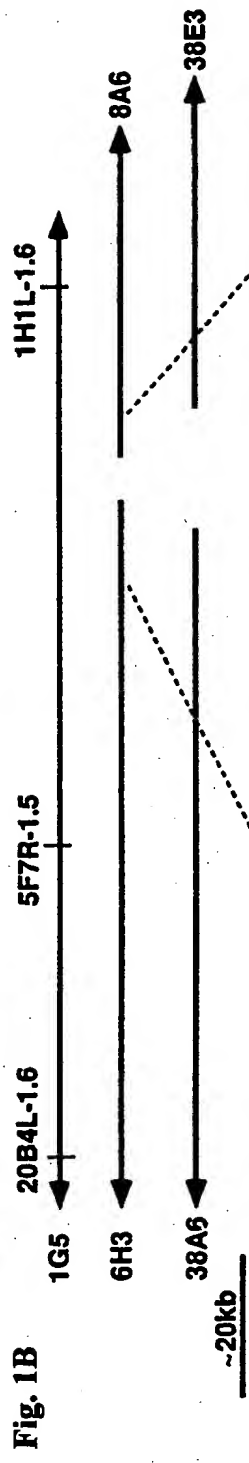
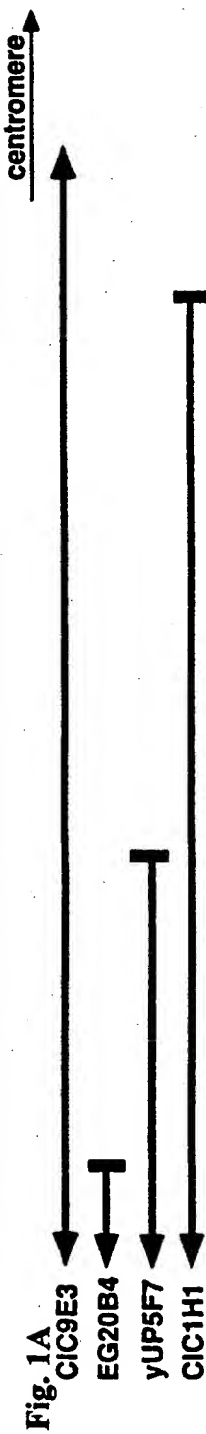
1. An isolated DNA sequence that encodes a LSD1 polypeptide.
2. The isolated DNA sequence of claim 1, wherein the sequence is selected from the group consisting of SEQ ID NO13, SEQ ID NO 14 and SEQ ID NO 15.
- 10 3. The isolated DNA sequence of claim 1, wherein the sequence comprises SEQ ID NO 13.
4. The isolated DNA sequence of claim 1, wherein the sequence comprises SEQ ID NO 15.
- 15 5. The isolated DNA sequence of claim 1, wherein the sequence comprises SEQ ID NO 15.
- 20 6. The isolated DNA sequence of claim 1, wherein the DNA is cDNA.
7. The isolated DNA sequence of claim 1, wherein the DNA is genomic.
- 25 8. The isolated DNA sequence of claim 1, wherein the polypeptide comprises SEQ ID NO 16.
9. The isolated DNA sequence of claim 1, wherein the polypeptide comprises SEQ ID NO 17.
- 30 10. A protein encoded by the isolated DNA sequence of claim1.
11. A chimeric construction comprising a promoter sequence and a DNA sequence according to claim 1.
- 35 12. A transformation vector comprising the isolated DNA sequence of claim 1.
13. A mutated DNA sequence derived from the DNA sequence of claim 1.

14. A transgenic plant expressing *LSD1* mutant genes that affect resistance to herbicidal compounds that normally result in plant cell death.
- 5 15. A transgenic plant expressing *LSD1* mutant genes which affect resistance to plant pathogens that normally result in plant cell death.
16. A messenger RNA encoding *LSD1*.
- 10 17. An isolated DNA sequence that encodes the zinc finger consensus selected from the group consisting of SEQ ID NOS 1-3.
18. A protein containing a zinc finger protein selected from the group consisting of CxxCxRxxLMYxxGASxVxCxxC, CxxCRxxLMYxxGASxRxVxCxxC,
15 CxxCxxLLMYxxGAxSxCxxC, CxxCxxLLxYxxGxxxVxCSSC,
CSGCRNLLMYPVGATSVCCAVC, CGGCHTLIMYIRGATSVQCSCC,
CGNCMMLLMYQYGARSVKCAVC, CGSCRRLLSYLRGSKHVKCSSC, and
CNNCKLLLMPYGYAPAVRCSSC, wherein x is any substituted amino acid.
- 20 19. A gene encoding a zinc finger protein according to claim 18.
20. An isolated DNA sequence encoding a protein according to claim 18.
21. A recombinant plant transformed with the DNA sequence as claimed in claim 1.
- 25 22. A recombinant plant transformed with the DNA sequence as claimed in claim 20.
23. An isolated DNA molecule that hybridizes under hybridization conditions to a DNA sequence as claimed in claim 1.
- 30 24. An isolated DNA molecule that hybridizes under hybridization conditions to a DNA sequence as claimed in claim 20.
25. An isolated DNA sequence that encodes a *LSD1* homologue.
- 35 26. The isolated DNA sequence of claim 25, wherein the homologue is selected from the group consisting of *LOL1* and *LOL2*.

27. The isolated DNA sequence of claim 25, wherein the homologue is selected from the group consisting of SEQ ID NO:48, SEQ ID NO:55, SEQ ID NO:60 and SEQ ID NO:62.
- 5
28. The isolated DNA sequence of claim 25, wherein the sequence is selected from the group consisting of SEQ ID NO:47, SEQ ID NO:54, and SEQ ID NO:59.
29. The isolated DNA sequence of claim 25, wherein the sequence comprises SEQ ID NO
10 47.
30. The isolated DNA sequence of claim 25, wherein the sequence comprises SEQ ID NO
54.
- 15 31. The isolated DNA sequence of claim 25, wherein the sequence comprises SEQ ID NO
59.
32. The isolated DNA sequence of claim 25, wherein the DNA is cDNA.
- 20 33. The isolated DNA sequence of claim 25, wherein the DNA is genomic.
34. A recombinant plant transformed with the DNA sequence as claimed in claim 25.
- 25 35. An isolated DNA molecule that hybridizes under hybridization conditions to a DNA
sequence as claimed in claim 25.
36. A protein encoded by the isolated DNA sequence of claim 25.
- 30 37. A chimeric construction comprising a promoter sequence and a DNA sequence
according to claim 25.
38. A transformation vector comprising the isolated DNA sequence of claim 25.
39. A mutated DNA sequence derived from the DNA sequence of claim 25.
- 35
40. A transgenic plant expressing *LOL1* mutant genes that affect resistance to herbicidal
compounds that normally result in plant cell death.

41. A transgenic plant expressing *LOL1* mutant genes which affect resistance to plant pathogens that normally result in plant cell death.
- 5 42. A messenger RNA encoding *LOL1*.
43. A transgenic plant expressing *LOL2* mutant genes that affect resistance to herbicidal compounds that normally result in plant cell death.
- 10 44. A transgenic plant expressing *LOL2* mutant genes which affect resistance to plant pathogens that normally result in plant cell death.
45. A messenger RNA encoding *LOL2*.
- 15 46. A nucleic acid that interacts with LSD1, selected from the group consisting of the nucleic acid sequences set forth in SEQ ID NOS:66-91.
47. A protein encoded by a nucleic acid according to claim 46.

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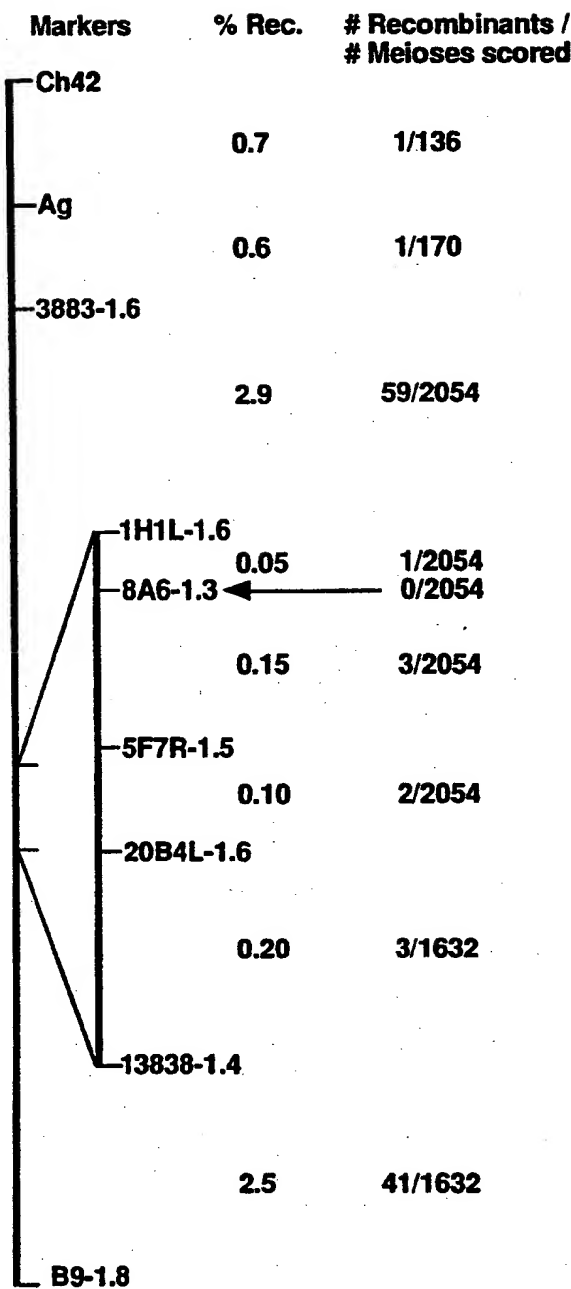


Figure 2

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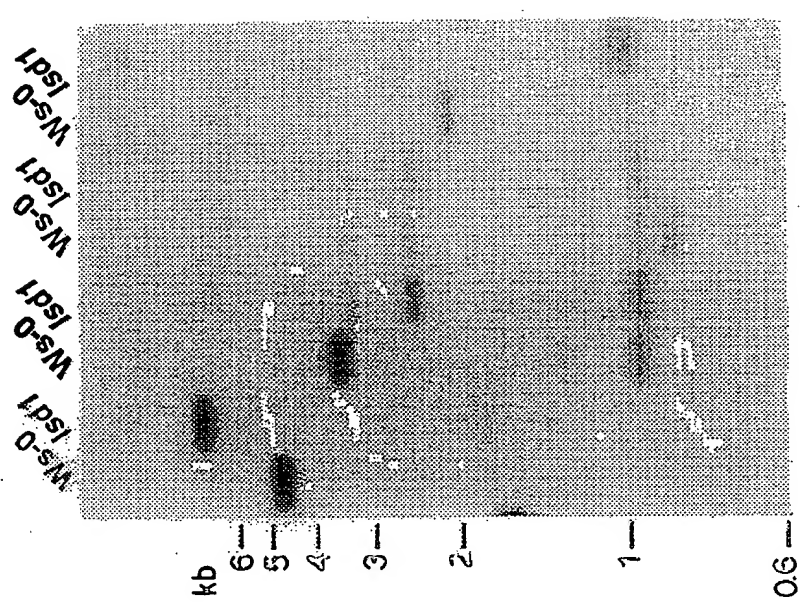


FIG. 3B

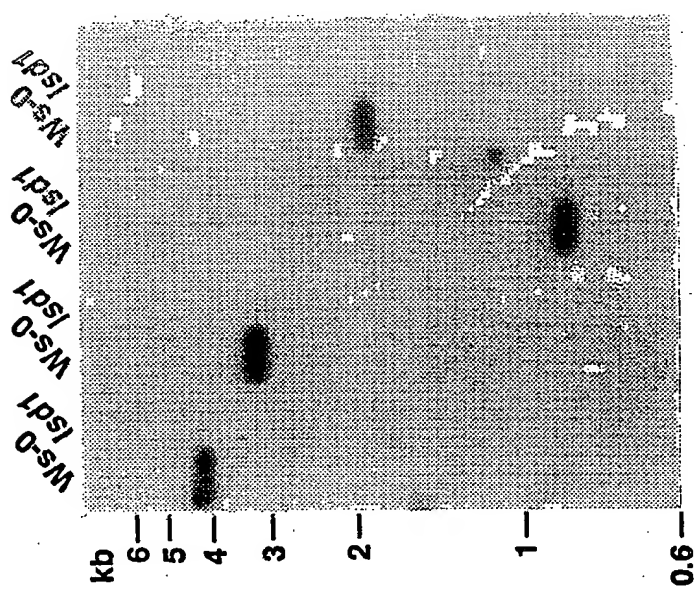


FIG. 3A

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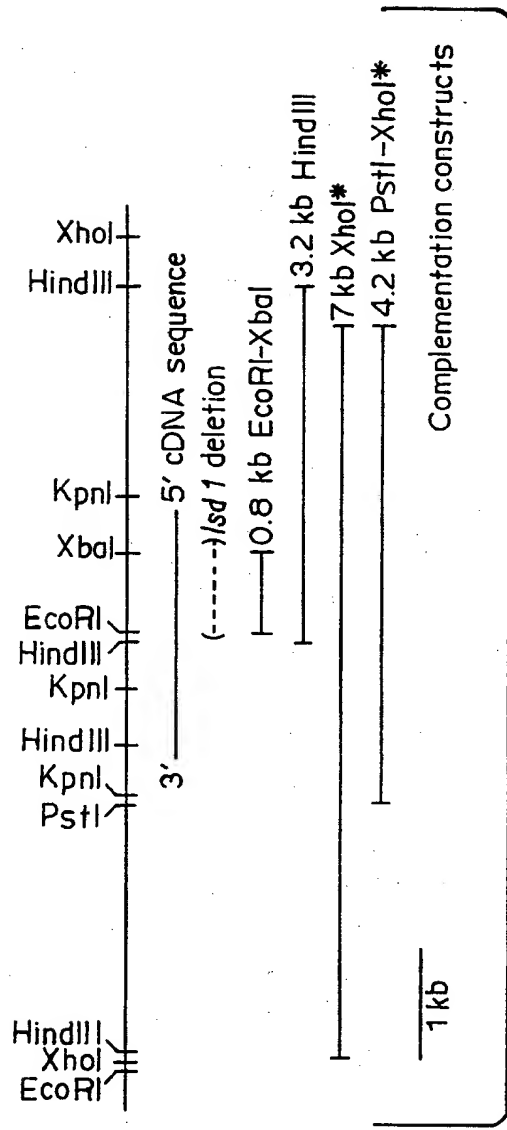


FIG. 3C

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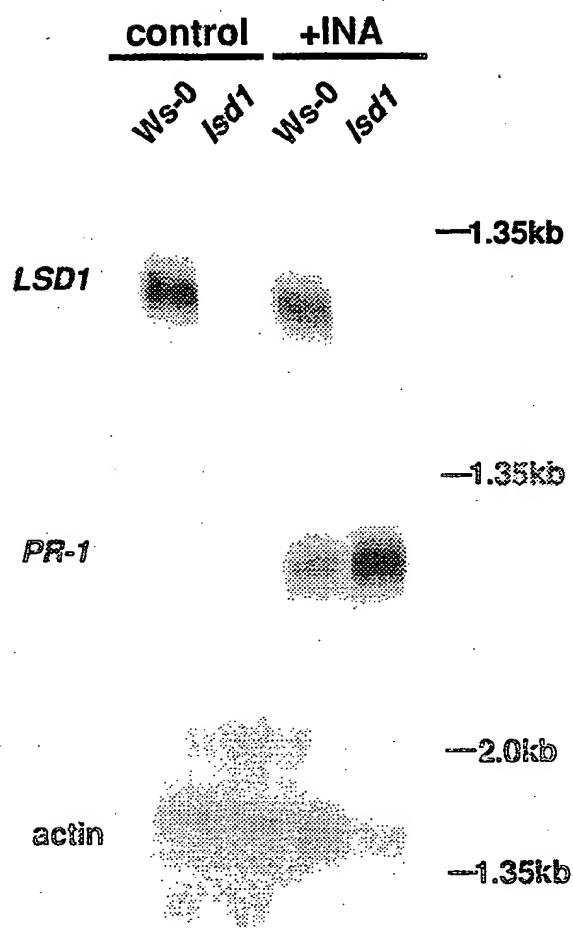


FIG. 4

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10 LVCHGCRNLIMYPRGASNVRCALCNTINMV 39
 ::| || :|| || || || || || || || || || ||
 51 IICGGGCRITMLMYTRGASSVRCCQFTNLV 80
 || || || || || || || || || || || || ||
 98 INCGHCRITLIMYPYGASSVKCAVCQFVTNV 127

Consensus:

L C CR IMY GAS V C C V
I + @

Figure 5

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Fig. 6A

Fig. 6B

Fig. 6C

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Fig. 7A

LSD1	10	LVCHGCRNLLMYPRGASNVRCALCNTINMV
	51	IICGGCRTMLMYTRGASSVRCSCCQTTNLV
	98	INCGHCRTTLMYPYGASSVKCAVCQFVTNV
consensus		C CR LMY GAS V C C V

Fig. 7B

LOL1	35	LVCSGCRNLLMYPVGATSVCCAVCNAVTAV
	74	LVCGGCHTLMLYIRGATSVQCSCCHTVNLA
	112	VNCGNCMMLLYQYGARSVKCAVCNFVTSV
consensus		C C LILMY GA SV C C V

Fig. 7C

LOL2	61	MVCGSCRLLSYLRGSKHVKCSSCQTVNLV
	99	VNCNNCKLLMYPYGAPAVRCSSCNSVTDI
consensus		C C LL Y G V CSSC V

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Fig. 8A

First zinc finger

LSD1

LVCHGCRNLLMYPRGASNVRCALCNTINMV

LOL1

LVCSGCRNLLMYPVGATSVCCAVCNAVTAV

LOL2

MVCGSCRRLLSYLRGSKHVKCSSCQTVNLV

consensus

VC CR LL Y G V C C V

Fig. 8B

Second zinc finger

LSD1

IICGGCRTMLMYTRGASSVRCSGCCQTTNLV

LOL1

LVCGGCHTLIMYIRGATSVQCSCHTVNL

LOL2

VNCNNCKLLMYPYGA
PAVRCSSCNSVTDI

consensus

C C I MY GA V CS C

Fig. 8C

Third zinc finger

LSD1

INCIGHCRTTILMYPYGASSVKCAVCQFVTNV

LOL1

VNCGNCMMLIMYQYGARSVKCAVCNFVTSV

consensus

NCG C I MY YGA SVKCAVC FVT V

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US98/04077

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) :Please See Extra Sheet.

US CL :435/320.1; 530/300, 324, 325, 326, 350; 536/23.6; 800/205

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/320.1; 530/300, 324, 325, 326, 350; 536/23.6; 800/205

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

Please See Extra Sheet.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X, P	DIETRICH et al. A Novel Zinc Finger Protein Is Encoded by the Arabidopsis LSD1 Gene and Functions as a Negative Regulator of Plant Cell Death. Cell. 07 March 1997, Vol. 88, pages 685-694, see entire document.	1-24
A	YANAGISAWA S. A novel DNA-binding domain that may form a single zinc finger motif. Nucleic Acids Research. 11 September 1995, Vol. 23, No. 17, pages 3403-3410, see entire document.	1-47
A	PUTTERILL et al. The CONSTANS Gene of Arabidopsis Promotes Flowering and Encodes a Protein Showing Similarities to Zinc Finger Transcription Factors. Cell. 24 March 1995, Vol. 80, pages 847-857, see entire document.	1-47



Further documents are listed in the continuation of Box C.



See patent family annex.

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P document published prior to the international filing date but later than the priority date claimed	

Date of the actual completion of the international search

06 MAY 1998

Date of mailing of the international search report

23 JUN 1998

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INTERNATIONAL SEARCH REPORT

International application No.
PCT/US98/04077

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	LIPPUNER et al. Two Classes of Plant cDNA Clones Differentially Complement Yeast Clacineurin Mutants and Increase Salt Tolerance of Wild-type Yeast. The Journal of Biological Chemistry. 31 May 1996, Vol. 271, No. 22, pages 12859-12866, see entire document.	1-47

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US98/04077

A. CLASSIFICATION OF SUBJECT MATTER:

IPC (6):

A01H 5/00, 7/00, 9/00, 11/00; C07K 7/08, 14/415; C12N 15/29, 15/63

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, DIALOG

search terms: LSD1, LOL1, LOL2, zinc finger, lesions simulating disease resistance, transcription factor, plant, cell death, LSD one like.

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